Wind-US User's Guide*

The NPARC Alliance

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^{*}This document describes the use of Wind-US Version 2.0, and was created February 9, 2009. It was originally based on the 24 Jan 1997 version of the NASTD User's Guide, by R. H. Bush, M. Mani, T. R. Michal, and W. W. Romer of McDonnell Douglas Aerospace. Please send documentation suggestions/corrections to Charlie Towne at towne@grc.nasa.gov. Questions about Wind-US itself or the NPARC Alliance should be sent to nparc-support@arnold.af.mil, (931) 454-7885.

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1 Introduction

This manual describes the operation and use of Wind-US, a computational platform which may be used to numerically solve various sets of equations governing physical phenomena. Wind-US represents a merger of the capabilities of four CFD codes — NASTD (a structured grid flow solver developed at McDonnell Douglas, now part of Boeing), NPARC (the original NPARC Alliance structured grid flow solver), NXAIR (an AEDC structured grid code used primarily for store separation analysis), and ICAT (an unstructured grid flow solver developed at the Rockwell Science Center and Boeing). Currently, the code supports the solution of the Euler and Navier-Stokes equations of fluid mechanics, along with supporting equation sets governing turbulent and chemically reacting flows.

1.1 Mathematical Model

All terms are retained in the governing equations, including secondary flow, reversed flow convection, pressure gradients normal to a wall, streamwise diffusion, and unsteady flow. All heat transfer terms are retained. Several algebraic, one-equation, and two-equation turbulence models are available. Transition may be specified through the use of an external file. Modification of the effective heat transport coefficient due to turbulence is linked to the momentum diffusion coefficient by a turbulent Prandtl number, which is taken to be constant.

The fluid may be treated as a thermally and calorically perfect gas, a thermally perfect gas, equilibrium air, or a mixture undergoing a finite-rate chemical reaction. For an ideal gas, conventional values are given to the gas constant R and the ratio of specific heats γ , or they may be specified.

1.2 Geometry and Mesh Description

Wind-US uses externally generated computational grids. Therefore, all geometric input and capability depend on the grid generator. Wind-US has no geometric input. All analyses must be preceded by a grid generation run.

Wind-US uses multi-zone computational grids, and is capable of computing solutions on a wide variety of structured or unstructured grids. However, not all of the features of Wind-US are available for both types of grids. The individual keyword descriptions note when a specific capability is limited to structured or unstructured grids.

Because Wind-US is written to accommodate arbitrary grid topologies and boundary condition combinations, it may be used to obtain solutions about most of the geometric configurations for which a grid can be generated. The multi-zone approach makes it possible to decompose virtually any configuration into a number of manageable subregions, or zones. Zonal connectivity information is computed using a pre-processing code (either GMAN or MADCAP), and stored in the grid file used by Wind-US. During the course of a solution, Wind-US maintains continuity in flow properties across zone boundaries through a process known as zone coupling (Romer and Bush, 1993).

¹Wind-US is a product of the NPARC Alliance, a partnership between the NASA Glenn Research Center (GRC) and the USAF Arnold Engineering Development Center (AEDC) dedicated to the establishment of a national, applications-oriented flow simulation capability. The Boeing Company has also been closely associated with the Alliance since its inception, and represents the interests of the NPARC User's Association.

1.3 Numerical Technique

The solution is executed iteratively on the computational mesh. The flow equations are evaluated using second-order-accurate finite differences. The partial differential equations are modeled in their conservative form. Explicit terms are computed using either upwind or central differencing, and their order may be controlled through the use of keywords in the input data file. The implicit terms are computed using either an approximately factored or four-stage Runge-Kutta scheme, or they may be disabled altogether. A Global Newton iteration scheme is also available, and may be used for unsteady flows with large time scales or as a convergence acceleration technique for steady flows.

1.4 Coding

Wind-US is written almost entirely in ANSI Standard Fortran 90. The low-level library routines are generally written in Fortran 77 and/or in ANSI C. The production version of the code is known to run on a variety of systems, including Linux, Silicon Graphics, Hewlett-Packard, Sun, and Cray.

2 Tutorial

This section is intended primarily for new users to demonstrate the simulation process using Wind-US. More experienced users may find this section useful as a road map through the simulation process and to help demonstrate new features. The approaches presented here are by no means unique, and detailed information is excluded by design. The user is referred to later sections of this User's Guide for more detailed information on the various aspects of running Wind-US, and in particular to Section 10 for more in-depth discussions on the choices available for each input keyword.

The approach taken here is to discuss the flow simulation process using as an example a simple subsonic internal flow in a diverging duct. The various files discussed in this section may be downloaded from the Wind-US documentation WWW site, at http://www.grc.nasa.gov/www/winddocs/user/tutorial.html#tutorial:downloading.

While it is clearly impossible to demonstrate every option in Wind-US with a single application, the basic mechanics of using Wind-US are demonstrated with this case. Additional abbreviated examples are also provided in Section 10 for specific keywords. For details of the flow simulation process for more complex cases, please see the various example applications accessible from the Wind-US Validation home page at http://www.grc.nasa.gov/www/wind/valid/validation.html.

The solution process using any conventional time-marching Navier-Stokes code is basically the same, and may be divided into the following steps:

- Gather information
- Create the computational grid
- Define the input
- Run the code
- Monitor convergence
- Examine the results

The mechanics of doing each of these steps may vary from code to code, however. The following sections describe how these steps are typically accomplished when using Wind-US.

2.1 Gather Information

As for any project, the first step is to gather all of the information required to completely specify the problem to be analyzed. Of course, as the flow simulation process proceeds, missing information will become apparent. The required information can be divided into three major categories — geometry, flow conditions, and boundary conditions. It is also important to understand the ultimate goal of the simulation. For example, is an accurate drag prediction required? Or, is lift required, but only to within 5%? The answers to these types of questions will determine the detail of the input information required to provide the necessary level of detail and accuracy in the solution.

2.1.1 Geometry

The more geometric details that can be determined for the target application, the more likely the results will provide an accurate simulation of the flow field. This is not to say that all geometric components must be modeled. Resolving fine geometric details of a configuration requires more grid points, and, as a result, longer run times. The level of detail to which the geometry must be modeled depends on the type of results required and the acceptable turn-around time.

The geometry of the Test Case 4 example is shown in Figure 1. The duct is 8 inches long, and the entrance and exit heights are 1 inch and 2 inches, respectively.

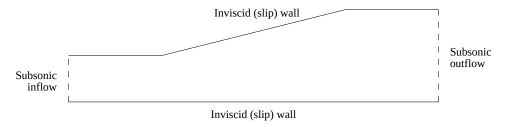


Figure 1: Test case geometry

The desired result from this calculation is the pressure distribution along the upper and lower walls, and the mass flow rate to within 10%. Thus, resolving the sharp surface gradients at the corners is not necessary. Other detailed geometric features such as weld joints, for example, are also not modeled. If the fine details of the boundary layer in the vicinity of the joints were important, then a significantly more detailed geometry description would be required.

2.1.2 Flow Conditions

In addition to the geometric information, flow conditions are also required, and are used to set the reference conditions used in non-dimensionalizing the governing equations solved by Wind-US. As with the geometric information, the simulation results will only be as good as the flow condition information provided by the user. Flow conditions should be specified that are representative of the flow being solved, so that the nondimensional variables used in the code are on the order of 1.0. A good choice is the inlet conditions for internal flows, and the freestream conditions for external flows.

For this example case, the inlet Mach number, total pressure, and total temperature are 0.78, 15 psi, and 600 °R, respectively. Starting from these conditions, the Reynolds number based on the duct height may be computed as 3.023×10^5 .

2.1.3 Boundary Conditions

Information is also required at boundaries that are at the "outer edges" of the computational domain. These boundary conditions are used to model the interaction between the flow inside the computational domain and surfaces or flows outside the domain. In fact, the boundary conditions are perhaps the most important factor influencing the accuracy of the flow computation.

Conditions at flow interface boundaries — boundaries between flows inside and outside the computational domain such as inflow, outflow, and freestream boundaries — must be known to the level of accuracy required by the simulation. For example, if flow rates are required to within 0.1%, even slight variations in total pressure at the inflow boundary must be specified. The number of conditions to be specified at a flow interface boundary depends on whether the flow is entering or leaving the computational domain, and whether it is subsonic or supersonic.

Information must also be specified at surface interface boundaries, such as solid walls and bleed regions. Simply specifying the type of boundary, such as an adiabatic no-slip wall, is often sufficient. Additional information may also be required, though, such as the wall temperature. The level of

detail that is needed for this information is determined, as discussed above, by the level of detail and accuracy required in the results.

Note that other types of boundaries may be present within the overall computational domain, that are not at the "outer edges." Multi-zone problems will have zonal interface boundaries. Some configurations will also have boundaries resulting from the grid topology, such as self-closing and singular axis boundaries. These types of boundaries need only be labeled.

For Test Case 4 the flow at both the inflow and outflow planes will be subsonic. Three conditions are needed at the inflow boundary, and one is needed at the outflow boundary. At the inflow boundary, uniform flow is specified, with total pressure and temperature equal to the inlet values of 15 psi and 600 °R. Since extreme accuracy in the solution is not needed, constant total conditions at the inflow are sufficient. At the outflow boundary, the exit static pressure is set to 14.13 psi. The Reynolds number for Test Case 4 is large enough that the boundary layers will have little influence on the pressure distribution within the duct. The upper and lower boundaries are therefore specified as inviscid (slip) walls.

In this tutorial, the procedure used to set boundary conditions when running Wind-US is discussed in Section 2.3.2. Additional details on all the boundary conditions available in Wind-US are presented in Section 5.

2.2 Create the Computational Grid

Wind-US uses externally-generated grids. The grids for all the zones must therefore be created before running Wind-US. The geometry of the application governs the overall shape of the boundaries, but the approach to gridding the flow field is not unique.

2.2.1 General Requirements

The Wind-US flow simulator provides considerable flexibility. Grid lines can conform to complex shapes or may pass through regions not in the flow field. The grid may be divided into zones to conform to the geometry better, to allow grid embedding (i.e., zones with finer grids in regions of high gradients like boundary layers), and/or to allow parallel computation. These zones may be abutting or overlapping, and overlapping grids may be single- or double-fringed.

Wind-US can compute flows using structured or unstructured grids. In this tutorial a structured grid is used. The indices (i,j,k) thus represent a curvilinear coordinate system, and physical Cartesian coordinates (x,y,z) are defined for each integer combination of indices. The handedness of both the physical and curvilinear coordinate systems is required to be the same at all points in the grid, i.e., both must be either left-handed or right-handed. Additionally, at least three grid points must fall between any two grid lines which represent a boundary within the computational domain. For example, if the k=1 boundary represents a solid surface and an adjacent k boundary represents a symmetry plane, the symmetry plane must be at k=5 or higher, so that the three points k=2,3, and 4 (at least) lie between the two boundaries.

The method used to create the grid is completely up to the user. For complex geometries, a sophisticated grid generation program is normally used. For very simple geometries, it may be easier to write a short program that constructs a grid using algebraic techniques.

2.2.2 Creating the Common Grid (.cgd) File

The computational grid used by Wind-US for a particular case is stored in a *Common Grid* (.cgd) file, so named because the file is formatted according to Boeing's Common File Format.² The (x, y, z) coordinates of all the grid points, zone coupling information, grid units, and scaling data are stored in this file.³

Since some grid generation codes do not produce .cgd files directly, a separate utility called cfcnvt is included with Wind-US that may be used to convert a variety of file formats, including PLOT3D files, to Common Files. A typical procedure is thus to first store the grid file as a PLOT3D xyz file, which is an available option in most general-purpose grid generation codes, and then convert it to a .cgd file using cfcnvt.

Zone coupling information is then added to the .cgd file using either the GMAN or MADCAP pre-processing utility. This is typically done at the same time as the boundary condition types are defined, and is discussed in Section 2.3.2. The .cgd file may then be examined to assess grid quality and list information about the points and zones in the grid. GMAN or MADCAP may also be used to generate the interior grid itself, given the grids on the zonal boundaries. Descriptions of these capabilities, and others, may be found in the user's guides for GMAN and MADCAP

2.2.3 Test Case 4 Grid

The computational grid for Test Case 4 was constructed by simple algebraic techniques using the following program, called *case4.mesh.f90*, which creates a PLOT3D xyz file in 2-D unformatted multi-zone form to Fortran unit 2, without an **iblank** array.

Program mesh

```
!----Purpose: This subroutine computes a 2-D three-zone grid for a
               simple diffuser, one of the test cases supplied with
ļ
               Wind-US. The grid is written to a file in PLOT3D 2-D
               unformatted form, without iblank'ing.
!----Called by:
!----Calls:
     Implicit none
!----Parameter statements
     Integer IDIM, JDIM
                           ! Max dimensions
     Integer NBLKS
                           ! Number of blocks
     Parameter (IDIM = 33, JDIM = 11)
     Parameter (NBLKS = 3)
!----Local variables:
     Integer i,j
                           ! Indices in x and y directions
     Integer iblk
                          ! Current block number
     Integer imax(NBLKS), jmax(NBLKS)
                                      ! Block grid sizes
     Real dx(NBLKS), dy(NBLKS)
                                     ! Non-dim grid increments in blks
```

 $^{^2}$ Wind-US also supports the use of CGNS files for the grid and flow solution, using the CGNSBASE keyword. This tutorial, however, uses common files.

³See the *Common File User's Guide* for details about the internal structure of common files.

```
Real x(IDIM, JDIM), y(IDIM, JDIM), z(IDIM, JDIM) ! Grid coordinates
     Real xdiff1,xdiff2 ! x at start/end of diffuser section
     Real xloc
                           ! Local x coordinate
     Real xmax
                          ! x at end of duct
     Real xstrt(NBLKS), xend(NBLKS) ! Non-dim x limits of blocks
                           ! Max y at x = xloc
     Real ymax
     Real ymax1,ymax2
                          ! Max y at start/end of diffuser section
     Real yslope
                           ! Slope of diffuser upper wall
     Real ystrt(NBLKS), yend(NBLKS)
                                    ! Non-dim y limits of blocks
!----Define geometric parameters
     Data xdiff1,xdiff2,xmax /2.0, 6.0, 8.0/
     Data ymax1, ymax2 /1.0, 2.0/
!----Set relative sizes and grid increments for each block
!----Block 1
     imax(1) = 17
     jmax(1) = 6
     xstrt(1) = 0.0
     ystrt(1) = 0.0
     xend (1) = 0.5
     yend (1) = 0.5
     dx(1) = (xend(1) - xstrt(1))/(imax(1) - 1)
     dy(1) = (yend(1) - ystrt(1))/(jmax(1) - 1)
!----Block 2
     imax(2) = 33
     jmax(2) = 11
     xstrt(2) = 0.0
     ystrt(2) = yend(1)
     xend(2) = xend(1)
     yend (2) = 1.0
     dx(2) = (xend(2) - xstrt(2))/(imax(2) - 1)
     dy(2) = (yend(2) - ystrt(2))/(jmax(2) - 1)
!----Block 3
     imax(3) = 17
     jmax(3) = 11
     xstrt(3) = xend(1)
     ystrt(3) = 0.0
     xend (3) = 1.0
     yend (3) = 1.0
     dx(3) = (xend(3) - xstrt(3))/(imax(3) - 1)
     dy(3) = (yend(3) - ystrt(3))/(jmax(3) - 1)
!----Open grid file, write header info
     Open (unit=2, file='fort.2', form='unformatted')
     Write (2) NBLKS
     Write (2) (imax(iblk), jmax(iblk), iblk=1, NBLKS)
```

```
!----Construct the grid
     yslope = (ymax2 - ymax1)/(xdiff2 - xdiff1) ! Slope of diff wall
     Do iblk = 1,NBLKS
                                  ! Loop over no. of blocks
                                  ! Loop over no. of x points
        Do i = 1,imax(iblk)
!----Compute x coordinates
           xloc = (xstrt(iblk) + dx(iblk)*(i-1))*xmax
           Do j = 1, jmax(iblk)
                                  ! Loop over no. of y points
              x(i,j) = xloc
           End do
!-----Get local max y, then compute y coordinates
           If (xloc <= xdiff1) then</pre>
                                              ! Upstream of diffuser
              ymax = ymax1
           Else if (xloc >= xdiff2) then
                                              ! Downstream of diffuser
              ymax = ymax2
                                              ! In diffuser
           Else
              ymax = ymax1 + yslope*(xloc-xdiff1)
           End if
           Do j = 1, jmax(iblk)
                                 ! Loop over no. of y points
              y(i,j) = (ystrt(iblk) + dy(iblk)*(j-1))*ymax
           End do
        End do
!----Write the grid file in PLOT3D xyz format
        Write (2) ((x(i,j),i=1,imax(iblk)),j=1,jmax(iblk)), &
                   ((y(i,j),i=1,imax(iblk)),j=1,jmax(iblk))
     End do
     Stop
     End
```

Figure 2 shows the resulting grid. Three zones are used, with grid sizes of 17×6 , 33×11 , and 17×11 , respectively.⁴

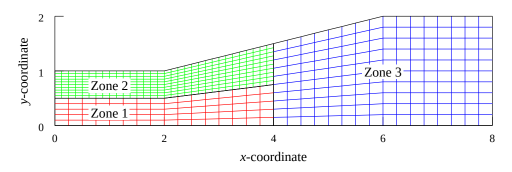


Figure 2: Test case grid

The grid file written by the above program, named case4.xyz, was converted to a .cgd file named case4.cgd using cfenvt, as shown in the following runstream. Lines in slanted type were typed by the user.

⁴This geometry is simple enough that a single-zone grid would be sufficient, but a three-zone grid is used for illustrative purposes.

```
% cfcnvt
 *********************************
* Warning: This software contains technical data whose export is
* restricted by the Arms Export Control Act (Title 22, U.S.C., Sec 2751,
* et seq.) or Executive Order 12470. Violation of these export-control
* laws is subject to severe criminal penalties. Dissemination of this
* software is controlled under DoD Directive 5230.25 and AFI 61-204.
 ************************************
         ***** Common File Convert Utilities *****
CFCNVT - Version 1.54 (last changed 2007/02/13 22:47:44)
  0: Exit program
  2: Import
            a Common File
  3: Compress a Common File
  4: Break Common File into multiple transfer files
  5: Combine multiple transfer files into Common File
  6: Append one Common File to another
  7: Convert Common File binary to a text file
  8: Convert Common File text to a binary file
 11: Convert PLOT3D/Pegsus file to Common File
 12: Convert GASP
                    file to Common File
 13: Convert OVERFLOW file to Common File
 14: Convert Common File to OVERFLOW file
 15: Convert CFPOST GPU file to Common File GPC
 16: Convert ascii rake to Common File rake CGF
 17: Convert Pegsus 4.0 files to Common File
 18: Convert Common File CFL to Plot3d Q
 Enter the number from one of the above requests
11
  PLOT3D file type menu
  0: Main menu
  1: Convert a PLOT3D Grid
                            (.x) file to CFS.
  2: Convert a PLOT3D Solution (.q) file to CFS.
  3: Convert a PLOT3D Function (.f) file to CFS.
 Enter the number from one of the above requests
  PLOT3D Number of Grids menu
  0: Main menu
  1: PLOT3D Single zone format.
  2: PLOT3D Multi zone format.
 Enter the number from one of the above requests
  PLOT3D Zone dimension menu
```

```
0: Main menu
  1: PLOT3D 2d zone format.
  2: PLOT3D 3d zone format.
 Enter the number from one of the above requests
  PLOT3D Format menu
  0: Main menu
  1: PLOT3D Formatted (ASCII).
  2: PLOT3D Unformatted (sequential binary).
  3: PLOT3D Binary (c binary).
 Enter the number from one of the above requests
  PLOT3D Iblank menu
  0: Main menu
  1: PLOT3D grid with IBLANK format.
  2: PLOT3D grid without IBLANK format.
 Enter the number from one of the above requests
  PLOT3D Precision menu
  0: Main menu
  1: PLOT3D Single precision format.
  2: PLOT3D Double precision format.
 Enter the number from one of the above requests
  PLOT3D INTOUT menu
  0: Main menu
  1: No INTOUT/XINTOUT file.
  2: INTOUT file.
  3: XINTOUT file.
 Enter the number from one of the above requests
 Enter PLOT3D .x file to convert with suffix
case 4.xyz
 Enter output Common File name with suffix
case4.cqd
                          1 17
2 33
 zone, imax,jmax,kmax:
                                                        6
 zone, imax,jmax,kmax:
                                                       11
```

```
33
                                                          11
 zone, imax,jmax,kmax:
                                                                        1
 Global maximums set as follows:
 mimax 33
 mjmax 11
 mkmax 1
 mpts: 363
 recl: 726
 Processing zone ZONE
 Writing mesh data
                                         6
                                                     1
 imax,jmax,kmax:
                           17
Processing zone ZONE
Writing mesh data
 imax,jmax,kmax:
                           33
                                        11
                                                     1
Processing zone ZONE
                        3
Writing mesh data
 imax,jmax,kmax:
                           17
                                        11
                                                     1
  0: Exit program
  2: Import
              a Common File
  3: Compress a Common File
  4: Break Common File into multiple transfer files
  5: Combine multiple transfer files into Common File
  6: Append one Common File to another
  7: Convert Common File binary to a text file
  8: Convert Common File text to a binary file
 11: Convert PLOT3D/Pegsus file to Common File
 12: Convert GASP
                       file to Common File
 13: Convert OVERFLOW file to Common File
 14: Convert Common File to OVERFLOW file
 15: Convert CFPOST GPU file to Common File GPC
 16: Convert ascii rake to Common File rake CGF
 17: Convert Pegsus 4.0 files to Common File
 18: Convert Common File CFL to Plot3d Q
 Enter the number from one of the above requests
0
```

2.3 Define the Input

The next step is to define the input data. Input is required to specify the flow and initial conditions, the boundary conditions, and various parameters controlling the physical and numerical models to be used when running the code.

The primary file controlling how Wind-US is executed is the *Input Data* (.dat) file. With many CFD codes the input data are specified using Fortran namelist and/or formatted input. With Wind-US, the input is specified using descriptive keywords. The formatting rules for the .dat file are described in Section 7.1.

This section is intended as an introduction to some of the more commonly-used keywords. After reading the information presented here, a new user should supplement it with the detailed information in Section 10. For many cases, the default values for the various keyword options are acceptable,

but users should become familiar with all of the options for the most effective use of Wind-US.

2.3.1 Descriptive Header and Comments

The first three lines of the file are reserved for geometry, flow condition, and arbitrary titles, respectively. Each of these titles may be up to 64 characters long.

Comment lines, beginning with a /, may be placed anywhere in the file after the first three lines. The readability of the .dat file may be improved significantly through the liberal use of comments — for example, to separate logical sections of the data file like boundary conditions, numerical operators, convergence monitoring parameters, etc.

2.3.2 Boundary Conditions

With most CFD codes, boundary conditions are completely specified in the input data file. With Wind-US, however, setting boundary conditions is a two-step process, defining first the *type* of boundary, and then any *values* that are needed.

The first step is to label each boundary of each zone with the *type* of boundary condition to use, such as "viscous wall," "outflow," or "coupled." This is done using either the GMAN or MADCAP pre-processing code, and the information is stored in the Common Grid (.cgd) file. Details on the boundary condition types available for use with Wind-US are in Section 5 and in the GMAN User's Guide.

Boundary condition types may be specified for all or part of a boundary, allowing multiple boundary conditions on a single boundary.

"Coupled" zonal interface boundaries do not have to be explicitly labeled by the user. Both GMAN and MADCAP can automatically examine the grid to find them and determine the zones involved, compute the geometric interpolation factors, and store the information in the .cgd file. GMAN or MADCAP is also used to cut holes and generate interpolation coefficients for overlapping ("chimera") boundaries. The process is currently not completely automated for chimera boundaries.

The second step is to define any *values* needed for a particular boundary condition, such as an exit pressure or a bleed rate. This information is specified in the *.dat* file. Keywords are available to specify conditions at inflow boundaries (ARBITRARY INFLOW), at outflow boundaries (COMPRESSOR FACE, DOWNSTREAM MACH, DOWNSTREAM PRESSURE, MASS FLOW), along solid walls (MOVING WALL, TTSPEC, WALL TEMPERATURE), in bleed and blowing regions (BLEED, BLOW), and across actuators and screens (ACTUATOR | SCREEN).

2.3.3 Flow and Initial Conditions

The flow conditions (Mach number, and static or total pressure and temperature, plus the angles of attack and yaw) are specified using the FREESTREAM keyword. These conditions, along with a reference length based on the units used in the .cgd file, are used as the reference conditions and determine the Reynolds number.

The usual procedure with Wind-US is to start a new problem by setting the initial conditions at each grid point equal to the values specified using the FREESTREAM keyword. Other keywords allow different values to be used in different zones (ARBITRARY INFLOW), a boundary layer to be added along a specified surface in a zone (BL_INIT), and reinitialization of the flow in specified zones after a restart (REINITIALIZE). Previously-run, partially-converged cases will normally be restarted using

the current solution as initial conditions (RESTART). More information about flowfield initialization may be found in Section 3.6.

2.3.4 Physical Model Controls

Dimensionality

Wind-US may be used for three-dimensional, two-dimensional, quasi-three-dimensional, or axisymmetric configurations. Internally, Wind-US treats the grid as three-dimensional, with indices i, j, and k. Two-dimensional cases simply have $k_{max} = 1$, with the i-j grid lying in a non-zero, constant z plane. The effect of area variation in an otherwise two-dimensional configuration may be modeled using Wind-US's quasi-three-dimensional capability, which is activated by setting the z-coordinate equal to the "width" of the geometry at each grid point. Axisymmetric configurations are modeled using a two-dimensional grid in conjunction with the AXISYMMETRIC keyword. More details may be found in Section 3.1.

Flow Equations

The default equations solved by Wind-US are the full Reynolds-averaged Navier-Stokes equations. Keywords are available to solve the Euler equations (TURBULENCE), parabolized Navier-Stokes equations (MARCHING), or the thin-layer Navier-Stokes equations (TSL). See Section 4.2.2 for more information about the thin-layer option.

Turbulence Model

A variety of turbulence models are available in Wind-US through the TURBULENCE keyword. These include the Baldwin-Lomax, Cebeci-Smith, and P. D. Thomas algebraic models; the Spalart-Allmaras and Baldwin-Barth one-equation models; and the Chien k- ϵ , Rumsey-Gatski k- ϵ algebraic Reynolds stress, and Menter Shear Stress Transport (SST) two-equation models.⁵ A laminar flow option is also available using the TURBULENCE keyword, and a laminar-turbulent transition region may be modeled using the TTSPEC keyword. See Section 3.3 for more information.

Gas Model and Chemistry

A variety of gas models are available in Wind-US to complete the equation set. The fluid may be treated as a thermally and calorically perfect gas, a thermally perfect gas (frozen chemistry), equilibrium air, or a mixture undergoing a finite rate chemical reaction. For a thermally and calorically perfect gas, the values of γ , the laminar and turbulent Prandtl and Schmidt numbers, and the gas constant may be set using the GAS, PRANDTL, and SCHMIDT keywords. Real gas chemistry models are selected using the CHEMISTRY keyword. Several different chemistry packages are available in the form of files containing thermodynamic data, finite rate coefficients, and transport property data, described in Section 7.11.

2.3.5 Numerical Model Controls

Time Stepping

In Wind-US, the number of iterations or time steps to perform in a given run is defined in terms of cycles and iterations per cycle. An iteration advances the solution one time step. A cycle consists of

⁵It should be noted that the algebraic models are older and infrequently used. They may not work as well as in some other codes, and there may be bugs in their implementation.

a solution pass over all the zones. Zone coupling, the process whereby Wind-US exchanges flowfield information between zones, only occurs at the end of each cycle. The Common Flow (.cfl) file is also updated only at the end of each cycle. The number of cycles to be performed is set using the CYCLES keyword, and the number of iterations per cycle, which may vary from zone to zone, is set using the ITERATIONS keyword. The default is five iterations per cycle.

The time step size is controlled by the CFL# keyword. By default, local time stepping is used, so that the solution advances at a different rate at each grid point. For unsteady problems, of course, the same time step size should be used throughout the flowfield, and an option is available with the CFL# keyword for this purpose. A Runge-Kutta time step formulation may be specified, using the STAGES keyword, and may be used for both steady and unsteady flows. Global Newton and second-order time marching are also available using options in the TEMPORAL keyword block.

See Section 4.1 for more information about cycles and iterations, and Section 4.5 for more about time step options.

Implicit Operator

The IMPLICIT keyword allows a variety of implicit operators to be specified, including point Jacobi, Gauss-Seidel, and MacCormack modified approximate factorization. Also available are options to: (1) turn off the implicit operator completely, resulting in an explicit calculation; (2) use a scalar (diagonalized) implicit operator; or (3) use a full block implicit operator. For these last three options, a different implicit operator may be specified for each computational direction.

The default is to use the full block operator in viscous directions, and the scalar (diagonalized) operator in inviscid directions.

The IMPLICIT BOUNDARY keyword may be used to specify that implicit boundary conditions are to be used on "wall" boundaries. This should improve stability when the CFL number is above about 1.3.

Explicit Operator

Through use of the RHS keyword, a wide variety of explicit operators are available for evaluation of the first-derivative terms on the right-hand side. These include a central difference scheme, the upwind Coakley, Roe, Van Leer, HLLE, HLLC, and Rusanov schemes, and modified versions of the upwind schemes (except Coakley) for stretched grids. Depending on the type of scheme used, the accuracy may be specified as anywhere from first to fifth order. The default is Roe's second-order upwind-biased flux-difference splitting algorithm, modified for stretched grids.

Damping Schemes

Various smoothing options are available in Wind-US to dampen instabilities that may occur under certain conditions. These include second- and fourth-order explicit smoothing (SMOOTHING, BOUNDARY-DAMP), and total variation diminishing (TVD) flux limiting for some of the explicit operators (TVD, BOUNDARY TVD). More details on the various damping options are in Section 4.3.

Convergence Acceleration

The ACCELERATE keyword may be used, in conjunction with the SMOOTHING and CFL# keywords, to increase the time step near the beginning of a calculation, in order to more quickly get through the start-up transients that may occur in the first few hundred iterations.

A grid sequencing capability is also available, using the SEQUENCE keyword, that may help speed convergence. With this option, grid points are removed from selected regions of the flowfield, re-

sulting in a coarse-grid solution which is obtained in a fraction of the time it would have taken for a full-grid solution. At the end of each run, the solution is interpolated back onto the original grid to aid in restarting the solution, and to provide a continuous flowfield for post-processing. The full grid should of course be used when the solution nears convergence.

See Section 4.6 for more information about the ACCELERATE option, and Section 4.2.1 for more about the grid sequencing capability.

Convergence Monitoring Parameters

The convergence criterion, in terms of the required value or reduction of the maximum residual, may be specified using the CONVERGE keyword. Integrated forces, moments, and/or mass flow may also be used to monitor convergence, by using the LOADS keyword to periodically compute and print these values for specified three-dimensional regions and/or two-dimensional areas of a computational surface.

For unsteady flow problems, where the time step is being specified in seconds and is the same throughout the flow field, a time history tracking capability is also available using the ${\tt HISTORY}$ keyword. Selected flow variables may be computed at specified grid points, and written to a separate ${\it Time \ History \ (.cth)}$ file.

More information about monitoring convergence is presented in Section 2.5 of this tutorial, and in Section 6.

2.3.6 Test Case 4 Input

Boundary Condition Types

GMAN was used in graphical mode to set the boundary condition types and store the information in the .cgd file. The interface boundaries between the three zones were automatically identified, and the geometric interpolation factors were computed and stored in the .cgd file. The inflow boundary (i=1 in zones 1 and 2) was labeled as "arbitrary inflow," the outflow boundary $(i=i_{max} \text{ in zone } 3)$ was labeled as "outflow," and the top and bottom boundaries (j=1 in zones 1 and 3), and $j=j_{max}$ in zones 2 and 3) were labeled as "inviscid wall."

The first step, obviously, is to start GMAN.

% gman

**** gman ****

Select the desired version from the following list.

- O) END
- 1) gman_pre optimized version

Single program automatically selected.

At this point, you may enter commands individually at the GMAN: prompt. Or, you could enter SWITCH or GRAPHICS to enter graphics mode.

The rest of this section describes in detail the use of GMAN for the tutorial test case. The graphics mode steps are on the left, with the Main Menu steps left-aligned and the Menu Options indented. Most of these are accomplished in GMAN by clicking on the listed menu item using the left mouse button. A few require entering text in the prompt area at the bottom of the screen. (See the "Graphical User Interface Basics" section of the *GMAN User's Guide* for a description of the various sections of the GMAN screen layout.)

The command line equivalents are shown on the right. Note that, in general, several graphics mode steps become consolidated into a single command.

We first need to tell GMAN the name of the file containing the grid.

Graphics Mode

Command Line Mode

 $\begin{array}{c} {\tt Common \ File} \\ {\tt \it enter} \ {\tt case4.cgd} \end{array}$

file case4.cgd

Next, we use GMAN's automated procedure to define zonal coupling information.

Graphics Mode

Command Line Mode

BOUNDARY COND.
AUTO COUPLE
RUN AUTO COUP

automatic couple face zone all

Next, for zone 1, we define the inflow and lower wall boundaries.

$\underline{\textbf{Graphics Mode}}$

PICK ZONE/BNDY

1: (from Zone List)

I1

MODIFY BNDY

CHANGE ALL

ARBITRARY INFLO

BOUNDARY COND.

YES - UPDATE FILE

PICK ZONE/BNDY

J1

MODIFY BNDY

CHANGE ALL

INVISCID WALL

BOUNDARY COND.

YES - UPDATE FILE

Command Line Mode

zone 1

boundary i1

arbitrary inflow

update

boundary j1

inviscid wall

update

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For zone 2, we define the inflow and upper wall boundaries.

Graphics Mode Command Line Mode

PICK ZONE/BNDY zone 2

2: (from Zone List)

I1 boundary i1 MODIFY BNDY arbitrary inflow

CHANGE ALL

ARBITRARY INFLO
BOUNDARY COND. update

YES - UPDATE FILE

PICK ZONE/BNDY

JMAX boundary jmax MODIFY BNDY inviscid wall

CHANGE ALL

INVISCID WALL

BOUNDARY COND. update

YES - UPDATE FILE

And for zone 3, we define the outflow and both wall boundaries.

Graphics Mode Command Line Mode

PICK ZONE/BNDY zone 3

3: (from Zone List)

IMAX boundary imax MODIFY BNDY outflow

CHANGE ALL OUTFLOW

BOUNDARY COND. update

YES - UPDATE FILE

PICK ZONE/BNDY

J1 boundary j1 MODIFY BNDY inviscid wall

CHANGE ALL

INVISCID WALL

BOUNDARY COND. update

YES - UPDATE FILE

PICK ZONE/BNDY

JMAX boundary jmax MODIFY BNDY inviscid wall

CHANGE ALL

INVISCID WALL

BOUNDARY COND. update

YES - UPDATE FILE

Finally, it's a good idea to check the boundary conditions to make sure all is OK.

Graphics Mode

Command Line Mode

TOP
CHECK zone all
CHECK BOUNDARY check boundary
PICK ZONE
ALL (from Zone List)
RUN BNDY CHKS

After hitting the Enter key to return to GRAPHICS mode, we can quit GMAN.

Graphics Mode

Command Line Mode

END exit

YES - TERMINATE

Input Data (.dat) File

The Input Data File for Test Case 4, named case 4.dat, is listed below. The explanatory notes, in italics, are not part of the file.

```
Wind-US test case 4, 2-D, 3 zones
                                             Titles
Subsonic internal flow
Run 1
/ Inlet conditions
Freestream total 0.78 15.0 600.0 0. 0.
                                             Inlet M, p_t (psi), T_t (°R), \alpha, \beta
/ Boundary conditions
Downstream pressure 14.13 zone 3
                                             Exit p (psi)
/ Numerics
Cycles 500
                                             Run 500 cycles
Iterations 5 Print frequency 5
                                             5 iterations/cycle; print every 5th
/ Viscous terms
Turbulence euler
                                             Solve inviscid equations
/ Convergence data
Loads
                                             Compute \dot{m} every 5 iterations at:
   print planes frequency 5
   zone 1
      surface i 1 mass
                                                   zone 1 entrance
   zone 2
      surface i 1 mass
                                                   zone 2 entrance
   zone 3
      surface i 1
                                                   zone 3 entrance
                       mass
      surface i last mass
                                                   zone 3 exit
Endloads
```

2.4 Run the Code

2.4.1 The wind Script

Wind-US is invoked using a Unix script, called wind, which prompts for the executable to be used (since production and beta versions of Wind-US may both be available on a system), the names of the various input and output files (which should be entered without the three-letter suffix), and for the queue in which the job is to run. If a multi-processing control (.mpc) file is present with the same base name as the .dat file (see Section 2.4.2), it also issues a prompt to verify that the job is to be run in parallel mode. The script then links the files to the appropriate Fortran units, and either starts Wind-US interactively or submits the job to the specified "at" or "batch" queue. Details on the wind script are in Section 8.1.

There are a couple of very convenient features built into the *wind* script. The first allows a run to be stopped at (or more exactly, shortly after) a pre-determined time through the use of an *NDSTOP* file. This is useful when an overnight run must be stopped before morning, when the workstations being used will be needed for interactive work. The second allows the user to break a long run into "sub-runs," by writing a script called *wind_post* containing tasks to perform between each run. This is useful, for example, when the complete solution is to be saved at various time intervals in an unsteady problem. Details on the use of the *NDSTOP* file and the *wind_post* script are in Section 7.9 and Section 8.2, respectively.

2.4.2 Parallel Operation

When Wind-US is run in parallel mode, multiple systems connected via a network work together as though they were a single computer. These systems are typically workstation class machines and need not be all from the same vendor.

A master-worker approach is used. Grid zones are distributed from the master system to the worker systems for processing. (Note that the master may also be a worker.) Each zone is solved in parallel with other zones on other systems. The systems exchange boundary information at the end of each solution cycle to propagate information throughout the flowfield. If there are fewer workers than zones, a worker will be assigned another zone when it finishes its current assignment.

The user specifies the names of the participating worker systems via a multi-processing control (.mpc) file, which must have the same base name as the .dat file. The user must of course have accounts on the master and worker systems, and the master must be allowed to communicate with each worker, and vice versa, using remote shell commands, and without entering a password. (See Section 9.3.2.) This is all that is required to utilize the parallel processing capability of Wind-US. The PVM software needed for parallel operation, and Wind-US itself, will be copied from the master to temporary directories on the workers.

Additional details about running Wind-US in parallel mode may be found in Section 9.

2.4.3 Running Test Case 4

To run Test Case 4, simply issue the *wind* command, and respond to the prompts as appropriate. The following terminal session shows how the case was run as a serial batch job on a Unix workstation. Lines in slanted type were typed by the user.

% wind -runinplace

Current solver settings are: --Solver set to Wind --Solver test mode set to NODEBUG --Solver debugger set to DEFAULT --Solver run que set to PROMPT --Solver run in place mode is set to YES --Solver cluster mode set to NO --Solver multi-processor mode set to NO --Solver run directory set to PROMPT --Solver bin directory set to /usr/local/wind --Solver clean up only mode set to NO --Solver status wait time set to 10 second(s) Select the desired version 0: Exit wind 1: Wind-US 1.0 2: Wind-US 2.0 Enter number or name of executable.....[2]: Basic input data.....(*.dat): case4 Output data.....(*.lis, <CR>=case4): Mesh file.....(*.cgd, <CR>=case4): Flow data file.....(*.cfl, <CR>=case4): ********************* case4.cfl does not exist, a fresh start will be performed. ***************** Enter a queue number from the following list or <CR> for default: 1: REAL (interactive) 2: AT_QUE Queue name.....(<CR> for 1): 2Enter at que (a,b..).....(<CR> for default): Deferred start time.....(hhmm [day], <CR>=now): Set stop flag at.....(hhmm [day], <CR>=don't set): Version...... /usr/local/wind/LINUX32-GLIBC2.3/XEON/bin/Wind-US2.exe Input file name....: case4.dat Wind output to....: case4.lis Grid file name....: case4.cgd Flow file name....: case4.cfl Job run que type is...: AT_QUE

***** Solver Run Script ****

Press <cr> to submit job, another key (except space) and <cr> to abort: job 3023 at 2007-10-01 12:02

There are several points to note from this terminal session.

• This case was run using the *-runinplace* option to the *wind* script, which means that Wind-US will be run in the current directory, and that output files will be written in the current directory.

The default is to run in a different (i.e., remote) directory, and is intended primarily for use with NFS-mounted home directories. In that case, it's faster to write the output files into a scratch directory on the system used to run Wind-US, rather than into the NFS-mounted home directory. The output files are automatically copied to the current directory at the end of the job.

Note that the terminology here is unfortunately a bit confusing. With an NFS-mounted home directory, running remotely really means running on a system different from the one the home directory is on. The "remote" system may actually be the local system originally logged onto.

If this case were run without the *-runinplace* option, the user would be prompted to enter the root name of the remote run directory, as follows:

Note the remote directory is assumed to exist on remote host
Enter the remote run root directory...(<CR>> for /tmp):

The full name of the remote run directory will be rootname/userid/basename.scr, where rootname is your response to the above prompt, userid is your userid, and basename is the base name of your .dat file. The default of /tmp for the root name implies that, generally, the "remote" system is actually the one the user logged into. It also means that, if you aren't using NFS-mounted home directories, and you forget to add the -runinplace option, no real harm is done. The output will be created under /tmp, then copied to the current directory when the job finishes.

- The default for the base name of the .lis, .cgd, and .cfl files is the same as that entered for the .dat file, and the three-letter suffixes should not be entered.
- A "fresh start" is being done for this case, since the .cfl file does not exist. If the .cfl file exists, Wind-US will automatically restart from the existing flow field.
- If the specified *.lis* file already exists, the output for the current run will be appended to the existing file. This typically occurs when restarting from a previous solution. If this isn't what you want, change the name of the existing *.lis* file before issuing the *wind* command.
- To run the job interactively, choose the REAL queue. This is intended mostly for debugging short runs, or checking that a case will successfully start. The user will be asked if the output should be written to the screen instead of the .lis file.

The AT_QUE queue is used to run a batch job using the Unix at or batch command. If the response to the Deferred start time prompt is defaulted, the job will be started immediately using the batch command. Any other response will result in the at command being used to start the job at the specified time.⁶

If batch queueing software is installed on the system being used, additional choices may be listed when the user is prompted for a run queue.

 $^{^6}Do$ not type now in response to the prompt. That will result in an "at" job being submitted, and a "too late" error.

• As noted earlier, if a multi-processing control (.mpc) file is present, the user is also asked to verify that the job is to be run in parallel mode, as follows:

```
A multiprocessing control file exists.... Do you want to run in multi-processor mode (y/n, <CR>=y):
```

2.5 Monitor Convergence

For complex real-world applications, it is generally not feasible to expect a converged solution in a single run. The times required to achieve convergence are generally too long and problems may occur which could corrupt the solution. Thus, executing the code several times and restarting from the previous solution is often the best approach. If problems do occur, the input parameters can be adjusted without starting from scratch.

Monitoring and properly assessing convergence levels during a Wind-US run (as well as examining the flowfield itself, as discussed in Section 2.6) are thus crucial in obtaining meaningful, useful results. Wind-US users may track convergence by following residuals and/or integrated forces, moments, and mass flow via the LOADS keyword. For engineering applications, the recommended convergence monitoring method is the tracking of integrated quantities of interest. For example, if a wing/body geometry is being modeled to determine drag, the integrated drag should be monitored and some reasonable bounds on drag oscillations should be used as the convergence criterion.

The solution residuals are included in the *List Output* (.lis) file. For each zone, Wind-US prints the zone number, cycle number, location of the maximum residual, equation number for which the maximum residual occurred, the value of the maximum residual, and the L₂-norm of all the residuals for all the equations over all the points in the zone. By default, the residuals are printed each iteration. The output interval may be changed, however, using the CYCLES and ITERATIONS keywords.

The integrated parameters that are chosen in the Input Data file via the LOADS keyword will also be listed in the .lis file. The integration may be done over a number of specified three-dimensional regions and/or two-dimensional areas of a computational surface.

For unsteady flow problems, where a constant time step is being specified and is the same throughout the flow field, a time history tracking capability may be used. Computed values of selected variables at specified grid points may be periodically written to a separate *Time History* (.cth) file. This capability is activated using the HISTORY keyword.

A utility included with Wind-US called *resplt* can be used to extract the residuals and/or integrated quantities from the *.lis* file, and create a GENPLOT file⁷ for post-processing.

An analogous utility called *thplt* can be used for the values stored in the .cth file.

Additional information about the various methods for monitoring convergence is presented in Section 6.

2.5.1 Test Case 4 Convergence

resplt was used to extract the maximum residual and the L_2 -norm of the residuals from the List Output (.lis) file and create GENPLOT files. As an example of the use of resplt, the following terminal session shows how a GENPLOT file containing the maximum residual was created. A

⁷The format of a GENPLOT file is described in the CFPOST User's Guide.

Wind-US User's Guide

GENPLOT file containing the L_2 -norm of the residual was created similarly, using selection "2". Lines in slanted type were typed by the user.

```
% resplt
                         **** resplt ****
        Select the desired version from the following list.
O) END
1) resplt optimized version
Single program automatically selected.
resplt - Version 1.10 (last changed 2007/02/14 17:14:15)
 * Warning: This software contains technical data whose export is
 * restricted by the Arms Export Control Act (Title 22, U.S.C., Sec 2751,
 * et seq.) or Executive Order 12470. Violation of these export-control
 * laws is subject to severe criminal penalties. Dissemination of this
 * software is controlled under DoD Directive 5230.25 and AFI 61-204.
 Enter full name of output list file:
case4.lis
Exit
                  0
Select Zone(s)
                 91
Select Frequency
                 92
Select average mode 99
Confined Outflow
 Mass Flow Ratio
 Back Pressure
                 16
 Average p0
                 93
Residuals
            Big
                L2
                         Integ. Planes Zone Grand
 NS
                 2
                           Force
                                  11
                                       5
            1
                          Lift
                                   17
 k-e
             3
                 4
                                       18
                                           19
 В-В
            20
                 21
                          Moment
                                  12
                                        6
                                            9
 S-A
            22
                 23
                          Momentum 13
                                       7
                                           10
 SST
             24
                 25
                           Mass 14
                                       26
NEWTON NS
            51
                 52
                           Heat Flx 54
                                       55
                 53
Time History
Enter Selection
 Reading residual data...
 1466 data points read from list file.
 Sorting residual data...
```

Sorting complete.

```
Enter FULL name of genplot file:
chist.big.gen
                       0
Exit
Select Zone(s)
                      91
Select Frequency
                      92
Select average mode 99
Confined Outflow
  Mass Flow Ratio
                      15
  Back Pressure
                      16
                      93
  Average p0
Residuals
                      L2
                               Integ. Planes Zone Grand
               Big
                      2
  NS
                 1
                                  Force
                                            11
                                                  5
                                                        8
                 3
                       4
                                                 18
                                                       19
  k-e
                                  Lift
                                            17
                                  Moment
  В-В
                20
                      21
                                                  6
                                                        9
                                            12
  S-A
                22
                      23
                                  Momentum 13
                                                  7
                                                       10
  SST
                24
                      25
                                  Mass
                                            14
                                                 26
NEWTON NS
                51
                      52
                                  Heat Flx 54
                                                 55
                      53
Time History
Enter Selection
0
```

The convergence history for Test Case 4, in terms of the maximum residual and the L_2 norm of the residual, is shown in Figure 3.

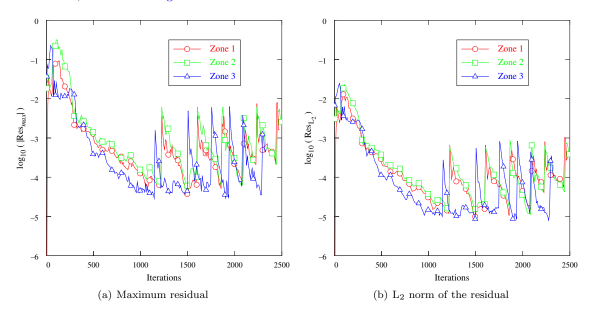


Figure 3: Test case convergence history in terms of residuals

For this case, the residuals decrease about three orders of magnitude in the first 1000 or so iterations, then oscillate about a relatively constant value for the remainder of the iterations. This behavior is not at all uncommon. As noted above, it's usually more meaningful to use a physical

quantity of interest when monitoring convergence. For this test case, one of the physical quantities of interest is the mass flow rate.

The Input Data file for this case specified that the integrated mass flux was to be computed at the entrances to all three zones, and at the exit of zone 3. GENPLOT files containing these integrated values were created using *resplt*, as illustrated above, using selection "14".

The time history for these parameters is shown in Figure 4, where the computed mass flux is plotted as a function of iteration number at the entrances to all three zones, and at the exit of the duct.

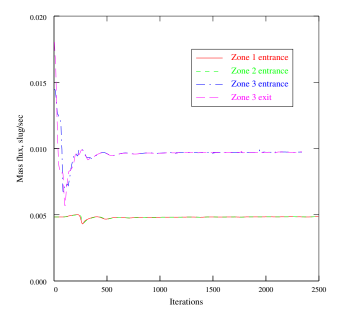


Figure 4: Test case convergence history in terms of mass flux

Based on the mass flux results, this case appears to converge within about 750–1000 iterations. Because this is a simple 2-D inviscid flow, with a coarse mesh, it converged quickly in a single run. The various convergence parameters were thus examined only at the end of the run. When running a more realistic, real-world configuration, convergence parameters like those shown in Figure 3 and Figure 4 should be examined at the end of each run. A more complete determination of convergence would also include examination of other physical quantities, such as the pressure distribution along the duct.

2.6 Examine the Results

Of course, the purpose of the solution process is to determine the features of the flow which can help answer the questions that drove the decision to perform the simulation in the first place. And, as indicated in the previous section, it's important to periodically examine the computed results during a run to help assess convergence and detect numerical problems that might be corrected by adjusting the input.

There are two types of information that can be extracted from the flow simulation — specific quantitative data and qualitative patterns. The first type includes things like pressure distributions,

drag, and total flow rate. The second type includes, for example, 2-D slices or full 3-D visualization of pressure contours, streamlines, or isothermal surfaces.

2.6.1 Using CFPOST

All flowfield results computed by Wind-US, including the mean flow variables, turbulence model variables, and chemistry variables, are written into a Common File called a *Common Flow* (.cfl) file.⁸ The CFPOST utility, included with the Wind-US distribution, is a post-processing tool for examining the contents of the .cfl file.

With CFPOST a wide variety of variables and integrated values may be computed. Listings of quantitative results may be sent to the screen or to a file. PLOT3D files may be created for other plotting packages and post-processors to use in displaying qualitative results. CFPOST can also be used to create x-y, contour, and vector plots directly, with PostScript output. Commands are available to precisely specify the information of interest, the domain of interest, and the units in which the results are to be presented. More details may be found in the *CFPOST User's Guide*.

2.6.2 Test Case 4 Results

The desired results from this calculation were the mass flow rate and the static pressure distribution. The mass flow rate is available in the List Output (.lis) file, as output generated via the LOADS keyword, and also from the GENPLOT files created while monitoring convergence. The result was 9.7×10^{-3} slug/sec.

To examine the static pressure distribution, CFPOST was first used to create a PLOT3D q file called *case4.q* from the Common Flow file *case4.cfl*, as follows:⁹

% cfpost

**** cfpost ****

Select the desired version from the following list.

- O) END
- 1) cfpost_prod optimized version

Single program automatically selected.

```
CFPOST_PROD - Version 4.6 (last changed 2007/08/24 22:35:29)
```

 $^{^8}$ As noted previously, Wind-US also supports the use of CGNS files for the grid and flow solution, using the CGNSBASE keyword.

 $^{^9}$ CFPOST can also be used to create a PLOT3D xyz file, but as noted in Section 2.2.3, an xyz file for this configuration was created during the mesh generation process.

```
1Wind-US test case 4, 2-D, 3 zones
Subsonic internal flow
```

Freestream conditions (* indicates calculated)

```
Mach number
                               0.780
Angle of attack
                               0.000 degrees
Yaw angle
                               0.000 degrees
Gamma
                                1.400
Gas constant (R)
                                                       1716.246 ft2/s2-R
                             287.000 m2/s2-K
Static pressure
                         69194.1
                                      N/m2
                                                    10.0357
                                                                lbf/in2
Static temperature
                         297.173
                                      K
                                                   534.912
                                                                R
Static density*
                        0.811292
                                      kg/m3
                                                  0.157417E-02 slug/ft3
                                                   15.0000
Total pressure*
                         103421.
                                      N/m2
                                                                lbf/in2
Total temperature*
                         333.333
                                      K
                                                    600.000
                                                                R
Total density*
                                                  0.209760E-02 slug/ft3
                         1.08106
                                     kg/m3
Speed of sound
                         345.549
                                      m/s
                                                   1133.69
                                                                ft/s
                                                                ft/s
Velocity
                         269.528
                                      m/s
                                                   884.280
Dynamic pressure
                         29468.4
                                      N/m2
                                                   4.27403
                                                                lbf/in2
Laminar Viscosity
                        0.183722E-04 \text{ kg/m-s}
                                                  0.383711E-06 slug/ft-s
Reynolds Number*
                        0.119020E+08 1/m
                                                  0.362774E+07 1/ft
Enthalpy*
                                      m2/s2
                         298511.
                                                  0.321314E+07 ft2/s2
                         334833.
                                      m2/s2
                                                  0.360412E+07 ft2/s2
Stagnation Enthalpy*
CFPOST> subset i all j all k all
CFPOST> zone 1
CFPOST> zone 2
CFPOST> zone 3
CFPOST> plot3d q case4.q 2d mgrid unformatted
Processing ZONE
                   1...
Processing ZONE
                   2...
Processing ZONE
                   3...
CFPOST> quit
```

The PLOT3D xyz and q files were then used as input to other post-processing packages to examine the computed results. The computed static pressure distribution along the upper and lower walls is shown in Figure 5, and the static pressure field is shown in Figure 6.

2.7 Summary

The steps in the generalized solution process listed earlier may be expanded and restated specifically for Wind-US as follows:

- Gather information detailed enough to specify the problem within the required accuracy, including the geometry, flow conditions, and boundary conditions.
- Create a grid file using any convenient grid generation software, saving the file in PLOT3D xyz format.
- Convert the PLOT3D xyz file to a Common Grid (.cgd) file using cfcnvt.
- Store the boundary condition types and zonal connectivity data in the .cgd file using GMAN or MADCAP.

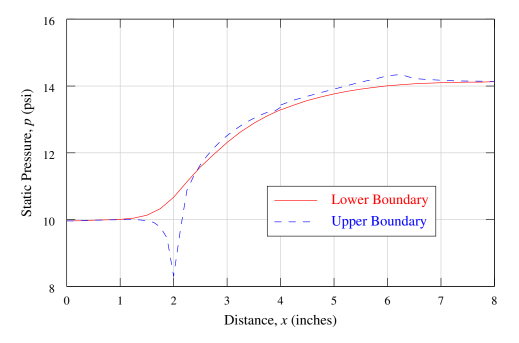


Figure 5: Test case static pressure distribution

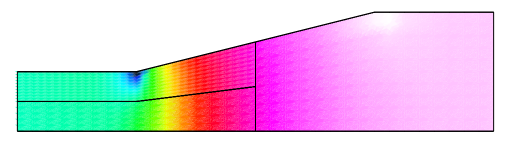


Figure 6: Test case static pressure field

- Prepare the Input Data (.dat) file, defining boundary condition values, initial conditions, program control parameters, and integrated parameters for monitoring convergence.
- \bullet For parallel execution, prepare the multi-processing control~(.mpc) file.
- Run Wind-US using the wind script supplied with the code.
- Monitor convergence by examining the residuals and integrated values in the List Output (.lis) file. The utilities *resplt* and CFPOST may be helpful.
- Periodically examine the computed results in the Common Flow (.cfl) file using CFPOST, creating PLOT3D files for other post-processing packages if desired.

3 Geometry and Flow Physics Modeling

3.1 Symmetry Considerations

Wind-US may be used with structured grids for axisymmetric, two-dimensional, or three-dimensional geometric configurations. Two-dimensional grids may be used not only for two-dimensional cases, but also for axisymmetric and area variation (quasi-three-dimensional) cases. Unstructured grids may be used only for three-dimensional configurations.

3.1.1 Three-Dimensional Cases

In three dimensions, each zone's computational mesh is comprised of six boundary faces and an interior grid. In structured grids, the mesh points are identified by three indices, usually labeled (i,j,k). In unstructured grids, each individual grid cell, and cell face, is numbered. Boundary conditions for each boundary face must be specified with GMAN or MADCAP before the grid may be used with Wind-US.

3.1.2 Two-Dimensional Cases

Two-dimensional cases may be run using structured grids only. The grid must be oriented such that the maximum k-index of the grid is one. In other words, a two-dimensional grid is defined by four boundary faces and an interior grid labeled by i- and j-indices. The grid must also reside in a non-zero, constant z-coordinate plane. The actual value used will not affect solution convergence or flowfield features, but it will affect flux-related post-processing calculations such as mass flow. For this reason, a value of 1.0 is recommended. Boundary conditions for the four boundary lines must be specified in GMAN or MADCAP.

3.1.3 Area Variation (Quasi-3D) Cases

With structured grids, the effect of area variation on two-dimensional computational models may be computed by using Wind-US's "quasi-three-dimensional" capability, which is activated simply through changes in the z-coordinate. The value of the z-coordinate is the "width" of the field at each grid point; the complete grid therefore represents the "width" variation of the field as a function of x and y. The important quantity to model is the ratio of cross-sectional areas between two adjacent axial stations. This means that the z-coordinate may be scaled with no effect on the computed flowfield, but a simple translation of the z-coordinates will change the computed flowfield, because the cross-sectional area ratio will be different. As with two-dimensional calculations, the value of the z-coordinate will affect flux-related post-processing calculations.

3.1.4 Axisymmetric Cases

Axisymmetric configurations may be modeled with structured grids, by using a two-dimensional grid generated at an arbitrary circumferential location on the geometry — e.g., the top centerline. Note that the grid should be generated on only one "side" of the configuration. Once again, the z-coordinate of the grid should be 1.0. The final step in using Wind-US's axisymmetric mode is the specification of the symmetry axis location and the circumferential sweep angle in the input data file. The circumferential sweep angle is the angle of the "pie shape" swept out by the grid about the

symmetry axis. Although the value of the sweep angle will not affect the computed flowfield, it will affect flux-related post-processing calculations.

Keywords: AXISYMMETRIC

3.2 Euler and Navier-Stokes Equations

Wind-US may be used to solve the Euler equations or the Reynolds-averaged form of the Navier-Stokes equations (Bush, 1988). All heat transfer and stress tensor terms are retained, and the equations are modeled in full conservation form. The effects of turbulence may be modeled using a variety of algebraic, one-equation, and two-equation turbulence models. Modification of the effective heat transport coefficient due to turbulence is linked to the momentum diffusion coefficient by a turbulent Prandtl number, which is assumed to be constant.

The fluid may be treated as a thermally and calorically perfect gas, a thermally perfect gas, equilibrium air, or a mixture undergoing a finite rate chemical reaction. For an ideal gas, conventional values are given to the gas constant R and the ratio of specific heats γ , or they may be specified. Effects of gravity (i.e., stratification) and rotation may also be included.

The equation set(s) to be solved must be specified in the input data file.

Keywords: TURBULENCE, GRAVITY, ROTATE

3.2.1 Freestream Conditions

Freestream flowfield conditions — Mach number, pressure, temperature, angle of attack, and angle of sideslip — must be specified in Wind-US's input data file. The Mach number must be greater than zero, and pressure and temperature may be specified as static or total values. These conditions are used to initialize the flowfield at the start of a run. For external flow problems, they are also applied at all inflow, outflow, and freestream boundaries during the course of a flow solution. For this reason, the outermost grid boundary should be far enough away from the body such that the freestream assumption is valid at the boundary.

Keywords: FREESTREAM

3.2.2 Reynolds Number Considerations

Wind-US does not allow the direct specification of a Reynolds number; freestream conditions must be specified that are consistent with the desired value. Currently, Wind-US uses a reference length which it computes based on the unit system of the grid file. The code prints out the reference length (in inches) that it uses, so that you may compare the Reynolds number from the code with the desired value. The Reynolds number in Wind-US's output is actually the Reynolds number "per reference length."

Here are a couple of examples.

Case I: Grid is same size as model

Suppose we have the grid for a wind tunnel model, and we want to run it at $M_{\infty} = 0.7$ and a Reynolds number of 12.1 million (based on the model length). We arbitrarily choose a total temperature of 520 °R (static temperature of 473.6 °R). Knowing the temperature, we can calculate

the speed of sound $(a = \sqrt{\gamma RT})$ and viscosity (from Sutherland's law). Using the Mach number, we can calculate the freestream velocity. Thus, we have

$$M_{\infty} = 0.7$$

 $T_{\infty} = 473.6$ °R
 $U_{\infty} = 746.67$ ft/sec
 $\mu_{\infty} = 3.474 \times 10^{-7}$ lb_f-sec/ft²
 $Re_L = 12, 100, 000$

We now have the Mach number and temperature for the Wind-US input data file, but we still need to calculate the freestream pressure. Using the definition of the Reynolds number (and the ideal gas law),

$$Re_L = \frac{\rho \, U \, L}{\mu} = \frac{P \, U \, L}{R \, T \, \mu}$$

or

$$P = \frac{RT \, \mu \, Re}{U \, L} = \frac{RT \, \mu \, Re_L}{U}$$

Note that Re_L is the Reynolds number per unit length of the physical model. For our example, if the model length is 10 inches,

$$Re_L = 12,100,000/(10/12 \text{ ft})$$

= 14,520,000(1/ft)

We can now calculate P:

$$P = \frac{RT \,\mu \,Re_L}{U} = \frac{(1716)(473.6)(3.474 \times 10^{-7})(14.52 \times 10^6)}{746.67}$$
$$= 5490.31 \, \text{lb}_f/\text{ft}^2 = 38.13 \, \text{psi}$$

We would now like to check our input. If we run Wind-US with the Mach, temperature and pressure specified above, the code will print a Reynolds number and a reference length near the top of the list output file. To obtain the desired Reynolds number, divide Wind-US's value of the Reynolds number by the output reference length and multiply by the model body length. This number may be compared with the desired model Reynolds number.

Case II: Grid is scaled from model size

Let us now assume that we want to run the previous grid at flight conditions, but we want to keep our same old 10-inch grid. We simply need to multiply the pressure by a scale factor. The equation now becomes:

$$P = \frac{RT \,\mu \, Re_L S}{U}$$

where

$$S = \frac{\text{Full Model Size}}{\text{Grid Model Size}}$$

For example, if we want to run a 100-inch wing using our 10-inch grid, S=10. If we want to run a flight Reynolds number of 26 million, we calculate P as:

$$\begin{split} P &= \frac{(1716.5)(473.6)(3.47\times10^{-7})(26\times10^6)(10)}{746.67} \\ &= 983113.0~\mathrm{lb_f/ft^2} = 682.72~\mathrm{psi} \end{split}$$

3.2.3 Mass Flow in Two-Dimensional Calculations

One of the options available in Wind-US is the specification of mass flow boundary conditions for subsonic duct analyses. Actual or corrected mass flow may be specified at duct exits, as may back pressure.

Within Wind-US and many post-processors, routines exist which integrate mass flow at desired computational planes. For 3D cases, the desired mass flow may be compared directly with the output from the integration routines:

$$\dot{m} = \int \rho u \, dA$$

However, for 2D calculations, the comparison is not so straightforward. There are three cases to consider.

Case I: 2D, Unit Depth

The first case involves running Wind-US on a truly two-dimensional grid of unit depth (z-coordinate is 1.0 everywhere). In this case, the input mass flow should be per unit depth. For example, let's say we want to run a 2D, unit depth model of a duct with a square exit. (If the exit were not square, this model would probably not be very good.) We would like a corrected mass flow of $500 \; \mathrm{lb_m/sec}$, and our actual model exit depth is 10 inches. If the grid input units are inches, we should ask for a mass flow of $50 \; \mathrm{lb_m/sec}$. If the grid input units are not inches, simply divide the actual mass flow by the z-coordinate value in inches.

Case II: 2D, Variable Width

When the z-coordinate is the width of the 2D grid, Wind-US adds in the area variation as a source to the 2D equations, making the analysis quasi-three-dimensional. In this case, the actual 3D mass flow should be specified in the input file. The integrated exit area will be (approximately, see Section 3.2.4 the real duct exit area, if the width has been specified correctly.

Case III: 2D, Axisymmetric

Axisymmetric runs require specification of the symmetry axis location and the circumferential angle subtended by the 2D grid. (This angle has no influence on the solution, but it does determine the area perpendicular to the grid.) The only reason this angle is an input parameter is so that you will know what the streamwise area is. In this case, the real exit geometry is circular, with a corresponding mass flow. The ratio of the input mass flow to the actual mass flow should equal the ratio of the input circumferential angle to 360. For example, if we are modeling a circular duct with a mass flow of 200 lb_m/sec using an axisymmetric model in Wind-US, and if we specify a circumferential angle of 36 (1/10 of 360), we should specify a mass flow of 20 lb_m/sec (1/10 of 200 lb_m/sec).

3.2.4 Mass Flow and Grid Areas

When dealing with subsonic duct analyses, you should be aware that the duct area as represented by the grid may be slightly different from the real area of the geometry being modeled, especially for ducts modeled with quasi-polar structured grids.

The duct area represented by the computational grid is often smaller than the real duct area, which, when running near critical mass flow, may prematurely choke the flow in the CFD solution. If the duct is circular and is modeled with a quasi-polar grid, the area error may be estimated.

Suppose we are modeling a circular duct with a quasi-polar grid using $k_{max} = 33$ circumferential points, each of which lie on the perimeter of the real duct at some streamwise station. If the circumferential points are evenly distributed, we may describe this topology as a circle which circumscribes a regular polygon of $k_{max} - 1 = 32$ sides. For a circle of radius R circumscribing a regular polygon of n sides, the area of the polygon is

$$A_p = \frac{1}{2}nR^2 \sin \frac{360}{n}$$

which means that the "grid area" is

$$A_g = \frac{1}{2}(k_{max} - 1)R^2 \sin \frac{360}{k_{max} - 1}$$

For our example, with $k_{max} = 33$, the grid area is 0.7% lower than the actual area.

There is no need to worry about this difference for most cases, but you should be aware of its possible effects.

3.2.5 Heat Transfer

At solid walls, Wind-US uses an adiabatic heat transfer boundary condition by default. A constant wall temperature may also be specified in the input data file.

Through the use of the TTSPEC keyword, point-by-point wall temperature distributions may also be specifed on boundary surfaces in structured grids. An auxiliary code, tmptrn, is used to create the wall temperature distribution, and write it into the common flow (.cfl) file.

The thermal conductivity is determined using a constant Prandtl number.

Keywords: WALL TEMPERATURE, TTSPEC

3.2.6 Viscosity

By default, Wind-US uses Sutherland's law to define laminar viscosity as a function of temperature. Keye's formula may be used in addition to Sutherland's law, and Wilke's law may be used to compute the laminar viscosity for multi-species flows.

Keywords: VISCOSITY

3.3 Turbulence Models

For turbulent calculations, Reynolds averaging assumptions are used to define a turbulent (eddy) viscosity, which is added to the laminar viscosity in the flow calculations. All the turbulence models available in Wind-US are coupled to the Navier-Stokes equations only through the turbulent viscosity.

For structured grids, users have a choice of several algebraic, one-equation, and two-equation turbulence models. In addition, various combined RANS/LES models may be used. For unstructured grids, two one-equation models and one two-equation model are available.

Note that a turbulence model (or inviscid or laminar flow) must be specified in the input data file. Wind-US will stop if you do not.

3.3.1 Algebraic Models

The algebraic turbulence models available in Wind-US for structured grids are the Cebeci-Smith model, the Baldwin-Lomax model (Baldwin and Lomax, 1978), and the P. D. Thomas model (Thomas, 1979), which adds a shear layer model to the Baldwin-Lomax model. After each iteration of the flow solver, these models compute the turbulent viscosity based on current flowfield quantities. Note that, because of their dependence on maxima and minima of flowfield variables, these models produce discontinuous turbulent viscosity distributions in the computed flowfield and require special numerical treatment at the juncture of two walls and on computational *i*-boundaries (the latter for historical reasons). The Baldwin-Lomax model is the most widely used algebraic turbulence model in Wind-US.

Keywords: TURBULENCE

3.3.2 One-Equation Models

Because of their efficiency and ability to produce continuous turbulent viscosity distributions, the one-equation turbulence models in Wind-US are the models of choice for many engineering applications. The one-equation models available in Wind-US for structured grids are the Baldwin-Barth (Baldwin and Barth, 1990) and Spalart-Allmaras (Spalart and Allmaras, 1992) models. The one-equation models available for unstructured grids are the Spalart-Allmaras and Goldberg pointwise models.

Keywords: TURBULENCE. FREE_ANUT

3.3.3 Two-Equation Models

Three two-equation turbulence models are currently available in Wind-US — the Menter Shear Stress Transport (SST) model (Menter, 1993; Mani, Ladd, Cain, and Bush, 1997), the low-Reynolds-number Chien k- ϵ model (Chien, 1982), and the low-Reynolds-number Rumsey-Gatski k- ϵ algebraic Reynolds stress model (Rumsey, Gatski, and Morrison, 1999; Rumsey and Gatski, 2000; Yoder, 2003).

The SST model may be used with both structured and unstructured grids. The two k- ϵ models are only available for structured grids.

The Menter Shear Stress Transport (SST) model is a blend of k- ϵ and k- ω . The equations are cast in k- ω form. Near the boundary the k- ω model is used, and the k- ϵ model is used away from the wall and in shear layers. Each equation is solved individually and an iterative method has been used on the left-hand side to reduce the factorization error. The model is robust, and may be more accurate in adverse pressure gradients than some of the other models in Wind-US. With structured grids, the SST model may be used with or without the compressibility corrections of Suzen and Hoffmann, and freestream values of k and ω may be specified.

The Chien k- ϵ model and the Rumsey-Gatski k- ϵ algebraic Reynolds stress model are also available in Wind-US. Several options may be specified with these models to control the initialization procedure, enhance stability, and improve accuracy in adverse pressure gradients and at high Mach numbers.

Keywords: TURBULENCE, COMPRESSIBLE DISSIPATION, PRESSURE DILATATION, FREE_K, FREE_OM, K-E ...

3.3.4 Combined RANS/LES Models

The idea behind combined RANS/LES (Reynolds-Averaged Navier Stokes / Large Eddy Simulation) turbulence models is to improve predictions of complex flows in a real-world engineering environment, by allowing the use of LES methods with grids typical of those used with traditional Reynolds Averaged Navier Stokes models. The combined model reduces to the standard RANS model in high mean shear regions (e.g., near viscous walls), where the grid is refined and has a large aspect ratio unsuitable for LES models. As the grid is traversed away from high mean shear regions, it typically becomes coarser and more isotropic, and the combined model smoothly transitions to an LES model.

Three different combined RANS/LES models are available in Wind-US for structured grids—the Spalart DES model, used in conjunction with the Spalart-Allmaras model (Spalart, Jou, Strelets, and Allmaras, 1997; Shur, Spalart, Strelets, and Travin, 1999), the SST model, with a limiter on ϵ , and the hybrid model of Nichols and Nelson, which may be used with both the SST and the Chien k- ϵ models (Nichols and Nelson, 2003).

The combined models may only be used for unsteady flows (i.e., the time step is a constant). They are zonal, however, so you can use a combined model in time-accurate mode in one zone, while using a standard RANS model in steady-state mode in the other zones.

Keywords: TURBULENCE, DES, LESB, HYBRID

3.3.5 Transition Specification

Through the use of the TTSPEC keyword, point-by-point transition data may be specified on viscous walls in structured grids. The data represent the percentage of turbulent viscosity to be added to the laminar viscosity at each grid point. An auxiliary code, *tmptrn*, may be used to create the transition data, and write it into the common flow (*.cfl*) file.

Keywords: TTSPEC

3.4 Gas Models

A variety of gas models are available in Wind-US to complete the equation set. The fluid may be treated as a thermally and calorically perfect gas, a thermally perfect gas (frozen chemistry), equilibrium air, or a mixture undergoing a finite-rate chemical reaction. Several different chemistry packages are available as files containing thermodynamic data, reaction rate data, and transport property data.

Keywords: CHEMISTRY

3.5 Other Models

For structured grids, various other models are available in Wind-US to model specific physical features of the geometry.

3.5.1 Actuator Disks

To simulate fan or compressor discontinuities, Wind-US provides an actuator disk modeling capability, which acts as a modification to the zone coupling boundary condition. The model assumes an infinitesimally thin disk and must be applied at a coupled zonal interface. The actual discontinuity is specified in the input data file as a solid-body rotation or free vortex flow, the effects of which are applied when transferring flow information between the two specified zone boundaries.

Keywords: ACTUATOR

3.5.2 Screens

Flowfield screens may also be modeled in Wind-US as discontinuities across coupled zonal interfaces. In the Wind-US input data file, one must specify the zones and boundaries between which the screen is located, the solidity of the screen, and one of several methods for calculating the losses through the screen. The screen model is not intended for use with choked screens, where the screen is significantly limiting the mass flow rate. During the solution start-up phase, it may be necessary to specify a low solidity, then increase it to the desired value to avoid strong choking in transients.

Keywords: SCREEN

3.5.3 Heat Exchangers

Heat exchangers may also be modeled at coupled zonal interfaces, using a procedure similar to that used for actuator disks and screens. The user specifies the zones and boundaries between which the heat exchanger is located, the temperature increase across the heat exchanger, and a static pressure loss coefficient.

Keywords: HEAT-EXCHANGER

3.5.4 Vortex Generators

Two models are available in Wind-US for including the effects of an array of vane-type vortex generators in three-dimensional flow. A vortex generator array consists of one or more vortex generators mounted on viscous wall boundaries.

The Wendt model (Dudek, 2006; Wendt, 2001) uses a discontinuous change in secondary velocity across a zonal interface boundary in order to simulate the vortices produced by the vortex generators. The model determines the strength of each vortex based on the generator chord length, height and angle of incidence with the incoming flow, as well as the incoming flow core velocity and boundary layer thickness.

The BAY model (Bender, Anderson, and Yagle, 1999) is a source term model which models the side force produced by the vortex generator and adds it to the momentum and energy equations. This side force automatically adjusts its strength based on the local flow. The user specifies the grid points over which the force is to be applied (i.e., enclosing each vortex generator).

Keywords: VORTEX GENERATOR

3.6 Flowfield Initialization

By default, Wind-US initializes the computational flowfield by setting the flow properties at each grid point equal to those specified with the FREESTREAM keyword in the input data file. The same initial conditions are applied at all points in the computational domain, including solid walls, zone boundaries, and freestream boundaries.

Several options in Wind-US lead to non-default initializations, including user-specified inflow conditions, boundary layer initialization, and reinitialization of portions of the flowfield on restart. The user-specified inflow and boundary layer initialization options are currently only available for structured grids.

3.6.1 User-Specified Initialization

Wind-US's ARBITRARY INFLOW keyword was designed to specify flow conditions at arbitrary inflow boundaries. However, it may also be used in a variety of ways to specify initial conditions in selected portions of the computational domain that are different from the freestream values. This capability may be useful, for example, when modeling a jet emanating from a solid wall. Zones downstream of the jet exit may have difficulty converging to the proper solution from conditions much different than the jet exit conditions.

To use the ARBITRARY INFLOW keyword as an initialization tool, specify the appropriate parameters and values for a zone as described below, and run Wind-US from scratch (i.e., without an existing flow (.cfl) file).¹¹

There are three ways that the ARBITRARY INFLOW keyword block may be used to modify the default initial conditions.

- If uniform inflow is specified for a particular zone using the UNIFORM keyword, the flow throughout that zone will be initialized to the conditions specified.
- The IJK_RANGE, XYZ_RANGE, and RTZ_RANGE keywords may be used to set flow conditions in specified regions. Up to 500 total regions are allowed (for unstructured grids, surfaces specified using USURFACE are included in this number). The default initial conditions will be used at points outside the specified region(s).
- The USERSPEC keyword may be used to specify a 1-D profile normal to the surface, translated through some buttline range, below the vehicle. Note that this option only applies to points in the i=1 computational plane. The default initial conditions will be used at the remaining points.

Keywords: ARBITRARY INFLOW

3.6.2 Boundary Layer Initialization

To provide a better approximation to near-wall flowfields, Wind-US provides a couple of boundary layer initialization options that may help speed the convergence of viscous flows near solid walls.

 $^{^{10}}$ Recall that the type of boundary, such as arbitrary inflow, is specified using GMAN or MADCAP, and stored in the common grid (.cgd) file, not in the input data file.

¹¹Note that since the ARBITRARY INFLOW keyword is also used to specify boundary conditions at arbitrary inflow boundaries, a conflict arises if (for some reason) the desired inflow properties are different from those being set as initial conditions. In this case, the ARBITRARY INFLOW keyword can still be used to set initial conditions by setting the number of cycles to be run to zero. Then, after changing the values specified with the ARBITRARY INFLOW keyword to the desired inflow values, simply restart using the initialized flowfield in the newly-created .cff file.

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First, by using multiple IJK_RANGE parameters with the ARBITRARY INFLOW keyword, and setting the range to include not only the inflow boundary but also locations downstream, one can set initial boundary layer profiles all along the viscous walls. Note that using this capability may add many lines to the input data (.dat) file. The INCLUDE keyword may be useful in keeping the main .dat file to a manageable size.

Another option is to use the BL_INIT keyword to specify a starting location and thickness for a laminar or turbulent boundary layer. However, this option may only be used on computational j-or k-boundaries, and only on one boundary in each zone.

Keywords: ARBITRARY INFLOW, BL_INIT

3.6.3 Reinitialization

In the event that portions of the computed flowfield become "polluted" with unrealistic flowfield data, due to numerical instabilities or other causes, you may wish to reinitialize portions of the flow. Wind-US's reinitialization option enables you to reset the flow conditions in specified zones.

For both structured and unstructured grids, conditions throughout the zone may be reinitialized to freestream values. In addition, for structured zones, conditions at all or specified grid points may be reinitialized to freestream values, or to values specified using the ARBITRARY INFLOW keyword and/or the BL_INIT keyword, as described above.

Keywords: REINITIALIZE, ARBITRARY INFLOW, BL_INIT

4 Numerical Modeling

Wind-US utilizes a number of different numerical methods in the solution of a selected set of field equations.

4.1 Iterations and Cycles

The usual method of obtaining a solution with Wind-US involves the initialization of the entire flowfield and the successive iteration of the flow solver, with appropriate boundary conditions, to a steady-state solution. Most boundary conditions are updated after each iteration of the flow solver; however, because of its expense, zone coupling may take place less frequently. Zone coupling is the process whereby Wind-US exchanges flowfield information between zones to maintain solution continuity at zonal interfaces. Updates of the flow file with the most current data also take place less frequently.

To facilitate flow file (see Section 7.3) updates and propagation of flowfield information to neighboring zones, Wind-US defines a "cycle" as a specific number of iterations performed in each of the zones, with zone coupling to effect information transfer at the end of each cycle. You must specify the number of solution cycles to be computed, and you may optionally specify the number of iterations per cycle to be calculated in each zone (or all zones simultaneously). By default, Wind-US performs five iterations per cycle in each structured zone, and one iteration per cycle in each unstructured zone.

For single-zone problems, you may wish to specify a large number of iterations per cycle, because there is no need for zone coupling operations; however, note that the flow file will be updated less frequently in this case. For multi-zone problems, a large number of iterations per cycle is not recommended, because it will greatly hinder the transfer of flowfield information between zones and, therefore, convergence.

4.1.1 "Turning Off" Zones

To "turn off" a zone in a multi-zone calculation, simply specify 0 or fewer iterations per cycle for that zone; values of -1 and -2 are used to indicate how zone coupling is to be handled for zones adjacent to the "dead" zone.

Keywords: CYCLES, ITERATIONS

4.2 Grid Considerations

Wind-US provides capabilities which allow you to make the most efficient use of the computational grid for a given configuration. The options described in this section will significantly reduce the CPU time required. However, the accuracy of the computations in one or more grid directions is also reduced, and they should therefore not be used haphazardly.

These options are only available for structured grids.

4.2.1 Grid Sequencing

Grid sequencing is a capability whereby Wind-US "removes" grid points from selected portions of the domain, resulting in a "coarse grid" solution which is obtained in a fraction of the time it would have taken for a "full grid" solution. For example, if you select one level of sequencing in all three directions in a particular zone, only every other grid point in each of the i-, j-, and k-directions will be used in the calculations, resulting in a decrease in CPU time of approximately 80–90%. (The speed-up is not a linear function of the number of sequencing levels.) Selecting two levels of sequencing would repeat the process, causing only every fourth grid point to be used. At the end of each run, the solution is interpolated back onto the entire grid to aid in restarting the solution and to provide a continuous field for post-processing.

Grid sequencing is very useful for converging gross flowfield properties before obtaining a final solution on the full grid. Note, however, that a sequenced solution may seem incorrect when post-processed using the entire computational grid, because of the interpolation process used at the end of each run. In addition, when restarting a full-grid solution from a sequenced solution, reducing the CFL number (time step) by 50% is highly recommended, in order to reduce the risk of solution instabilities which arise during the transition.

Keywords: CFL#, SEQUENCE

4.2.2 Thin-Shear-Layer Calculations

Another time-saving option available in Wind-US is the ability to solve the thin-shear-layer Navier-Stokes equations, instead of the full equations. By requesting that Wind-US remove the viscous terms from the equations in one or more computational directions, a CPU savings of up to 30% may be realized. This capability requires careful specification, because viscous terms must be retained along grid lines which are perpendicular to boundary layers, free shear layers, and other highly viscous flow features.

Because some solution instabilities have been linked to the use of the thin-shear-layer option, it is recommended that you disable this option near the completion (convergence) of your solution, creating a fully viscous flowfield model.

Keywords: TSL

4.3 Explicit Operator

The first-derivative (non-viscous) terms on the right-hand side of Wind-US's equation set are calculated by the code's explicit operator. You may choose from a wide variety of explicit operators, independently of the implicit (left-hand-side) operator. An explicit operator is selected by its type and accuracy.

For structured grids, the explicit operators available include a central difference operator, the upwind Coakley, Roe, Van Leer, HLLE, HLLC, and Rusanov schemes, plus modified versions of all these schemes (except Coakley) for stretched grids. Depending on the type of scheme used, the accuracy may be specified as anywhere from first to fifth order. The default is Roe's second-order upwind-biased flux-difference splitting algorithm, modified for stretched grids.

For unstructured grids, only the cell-centered Roe, HLLE, HLLC, and Rusanov operators are available. The accuracy may be first or second order.

Keywords: RHS

4.3.1 Explicit Smoothing

Certain types of explicit operators — for example, the central difference operator for structured grids — may require the addition of numerical smoothing in order to dampen instabilities which are a natural part of the scheme. Smoothing must also be added explicitly when utilizing Wind-US's convergence acceleration capability. Values for various smoothing parameters may be specified in the input data file for both structured and unstructured grids.

Keywords: ACCELERATE, SMOOTHING, LAX DAMPING

4.3.2 Explicit Boundary Damping

In flowfields dominated by acoustic phenomena, unwanted wave reflections may sometimes occur at freestream computational boundaries. To eliminate these unphysical reflections, Wind-US's boundary damping capability may be employed with structured grids to apply explicit smoothing near the freestream boundaries, with the effect of absorbing the reflected waves.

Keywords: BOUNDARY-DAMP

4.3.3 Total-Variation-Diminishing (TVD) and Slope Limiting

Various Total-Variation-Diminishing (TVD) limiters (for structured grids) and slope limiters (for unstructured grids) are available in Wind-US.

For structured grids, the TVD scheme limits local maxima and minima in flux quantities to prevent non-physical instabilities from arising during the solution. The default Roe scheme, for example, requires interpolation and/or extrapolation of flux quantities, which may produce minima or maxima that are outside the range of the original data. TVD operators attempt to limit these peaks to acceptable values based on the data used in the interpolation and extrapolation.

Three different TVD operators are available in Wind-US for structured grids, and may be specified on a zonal basis. For two of the operators, the severity of the limiting may be controlled through an input parameter.

Because only single data points are used in the first-order schemes, TVD, which limits interpolations and extrapolations, has no effect on such cases. It will only have an effect for second- and higher-order calculations. When attempting to model precise phenomena, such as acoustic waves, TVD should *not* be used, because it may produce undesirable damping or smearing of flowfield features. However, in most cases, TVD provides significant protection against numerical instabilities with less than a 10% increase in CPU time.

For unstructured grids, slope limiting is used by default for viscous cases, and for inviscid cases with second-order spatial differencing. Three procedures are available, the default standard slope limiter, a "super-bee" slope limiter, and Venkatakrishnan slope limiting. The severity of the limiting may be controlled through an input parameter.

Keywords: TVD, BOUNDARY TVD

4.4 Implicit Operator

The left-hand side of Wind-US's equation set is computed in the code's implicit operator, which may be set on a zonal basis.

For structured grids, options are available to select scalar or full block implicit operators, or to turn off the implicit operator completely for an entirely explicit calculation. With these options, a different implicit operator may be specified for each computational direction. By default, Wind-US uses the full block operator in viscous directions, and the scalar (diagonalized) matrix solver in inviscid directions.

A variety of other implicit operators are also available for structured grids, including point Jacobi, Gauss-Seidel, MacCormack's modified approximate factorization, and the ARC3D 3-factor diagonal scheme as used in the OVERFLOW code.

For unstructured grids, either a point or line Gauss-Seidel operator may be used, and the equations may be solved as either a full block or diagonal matrix system.

Keywords: IMPLICIT

4.4.1 Implicit Boundaries

In addition to the normal explicit boundary conditions, implicit boundary conditions may be used on "wall" boundaries in structured grids. These improve stability when the CFL number is above about 1.3.

Keywords: IMPLICIT BOUNDARY

4.5 Time Step

The time step can be one of the most important parameters in a Wind-US solution. It may mean the difference between a successful run and a code bomb. For steady flow problems, the time step is represented by the CFL (Courant-Friedrichs-Lewy) number, which is a non-dimensional time step based on the actual time step, the local grid spacing, and the local characteristic velocity (u + c). For unsteady flow problems, the time step in seconds must be specified directly.

4.5.1 CFL Number, Structured Grids

For stability reasons, the default CFL number is 1.3, which is sufficient for a wide variety of problems using structured grids. For inviscid solutions, the value may be raised slightly, while for highly complex flows (due to complex geometry or gas models), it may need to be lowered significantly. Often, in the beginning of a solution, or during the transition between a sequenced and fine grid solution, the CFL number should be set to 0.5 while transients settle out of the flowfield.

If you must consistently set the CFL number to 0.1 or lower to maintain stability in the solution, you should attempt to locate other possible causes of numerical instability, such as grid anomalies or grid/flow interactions. Such low CFL numbers translate into small time steps and unacceptable convergence rates.

When a CFL number above 1.3 is desired, the implicit boundary conditions should be used.

By default, Wind-US computes the time step from the input CFL number by using the minimum time step associated with the eigenvalues of each coordinate direction individually. That is:

$$dt = \min\left(\frac{\text{CFL}}{\lambda_x}, \frac{\text{CFL}}{\lambda_y}, \frac{\text{CFL}}{\lambda_z}\right)$$

Using TEST 105, the user may define the time step in various other ways, including the more conventional formula

 $dt = \frac{\text{CFL}}{\lambda_x + \lambda_y + \lambda_z}$

which provides a somewhat smaller effective time step.

Options in Wind-US allow the CFL number or time step to be gradually increased as the calculation proceeds, or to use the time step calculation procedure from the OVERFLOW CFD code.

For most calculations, the time step is computed based on a constant CFL throughout the field, which means that, for non-uniform grids, a local time step is used, and the solution advances at different rates in different parts of the grid. Because the CFL number is directly proportional to the physical time step and inversely proportional to the local grid spacing, the physical time step must be small where the grid spacing is small, in order to maintain a constant CFL number. For example, in boundary layers and shear layers, where the grid is often closely packed for better resolution, the solution advances more slowly than in other parts of the flowfield. In fact, most of the iterations in a viscous solution are spent converging viscous regions. In practice, the major flow features (e.g., shocks and other major pressure gradients) will develop quickly, while detailed flow phenomena will require a larger number of iterations.

Wind-US therefore uses by default a "cross-flow" CFL number which multiplies each coordinate direction's eigenvalues by a factor before determining the time step associated with that spacing and flow conditions. This factor varies from 1.0 when the flow is along the coordinate direction, to a user-specified value when the flow is perpendicular to the coordinate direction. The far-field CFL number is generally the value specified in the input data file with the CFL# keyword, but near the wall the code effectively uses a larger value. This should increase convergence of the boundary layer.

The cross-flow CFL factor is defined using the CROSSFLOW keyword. Values greater than one effectively increase the time step in the boundary layer (where the flow is parallel to the wall, but the minimum spacing is normal to the wall) by the specified factor. The default is 2.0 for three-dimensional flow, and 1.0 (i.e., no time step increase in the boundary layer) for two-dimensional flow and axisymmetric flow.

Keywords: CFL#, CROSSFLOW

4.5.2 CFL Number, Unstructured Grids

With unstructured grids in Wind-US, a larger CFL number is normally used than for structured grids. The default CFL number for unstructured grids is 10, and values as high as 50 may be used as the calculation proceeds.

Wind-US computes a local time step for each cell from the input CFL number by using the minimum time step associated with the eigenvalues at each face of the cell. That is:

$$dt = \min\left(\frac{\text{CFL}}{\lambda_{f1}}, \frac{\text{CFL}}{\lambda_{f2}}, \cdots\right)$$

Options in Wind-US allow the CFL number to be automatically increased or decreased as the calculation proceeds.

4.5.3 Runge-Kutta Time Step

For structured grids, several Runge-Kutta time step formulations are available for use in Wind-US. A traditional four-stage Jameson-type scheme or a three-stage, minimal-storage scheme may be selected in the input data file. You may also specify algorithm coefficients for each of the stages in the different schemes.

Runge-Kutta schemes attempt to provide a more accurate approximation to the solution at the next time level by advancing through several stages, or sub-iterations, to get there. In Wind-US, each sub-iteration may be one pass through the explicit operator with or without a pass through the implicit operator. (The original Runge-Kutta scheme was formulated using only an explicit operator.) Depending on the coefficients used, Runge-Kutta time step algorithms are often higher order in time than the default Euler time stepping algorithm. This makes them ideally suited for time-accurate calculations (see below).

Keywords: STAGES

4.5.4 Time-Accurate Solutions

The objective of time-accurate CFD calculations is the modeling of unsteady flow phenomena, such as vortex pairing or bluff body wakes. In unsteady calculations, a global physical time step must be used throughout the entire flowfield, regardless of the local grid distribution. Because the CFL number has a limit which approximates the upper limit of numerical stability, the global time step must be obtained from the maximum CFL number and the smallest grid spacing in the field. Thus, the stability of the calculations in regions of dense grid packing determines the time step for the entire flowfield.

The options specified with the CFL# keyword determine whether the input value is interpreted as a CFL number, or a physical time step. For single-zone time-accurate calculations with a structured grid, an option is also available to set a global time step, equal to the minimum time step in the zone based on the specified CFL number. However, for multi-zone time-accurate calculations, you must calculate the time step yourself, based on the desired CFL number and the minimum grid spacing, and specify the physical time step in the input data file.

For unsteady flows, the use of second-order time marching is recommended, possibly in combination with Global Newton iteration.

Keywords: CFL#, IMPLICIT ORDER, NEWTON, TEMPORAL

4.5.5 Global Newton Iteration

A Global Newton iteration technique is available in Wind-US for both structured and unstructured grids, and has proven to provide time-accurate solutions with large CFL numbers for unsteady flows with large time scales (Tramel and Nichols, 1997). The Global Newton technique stabilizes the solution and improves time accuracy by placing the entire unsteady transport equations on the right-hand side of the matrix solver, and iterating within a time step over all the zones. Thus, the interface boundaries are brought up to the new time level, along with the interior flow field, resulting in an essentially implicit treatment of the boundaries. The Global Newton algorithm has also been shown to improve steady-state convergence rates (Nichols and Tramel, 1997).

Keywords: NEWTON, TEMPORAL

4.6 Convergence Acceleration

The solution transients that occur in iterating from the freestream-initialized flowfield often cause the most severe stability problems in Wind-US. As the solution approaches convergence on a sequenced grid, most of these transients will have settled out of the field, with the change to fine grid being the only remaining "jolt" to the solution. Numerical stability may be forced through the addition of artificial numerical dissipation to the equations, but such dissipation may lead to non-physical results in the steady state solution.

The convergence acceleration capability in Wind-US allows the use of a large CFL number at the start of a calculation, with artificial dissipation used to eliminate stability problems, to get past the starting transients more quickly. The CFL number and the dissipation are then lowered as the calculation proceeds.

For structured grids, you may specify the starting and ending values of the CFL number, the initial amount of artificial dissipation (smoothing) to be added to the equations, the iteration numbers between which the smoothing should be applied and CFL number altered, and the zone(s) in which you would like to enable this capability. At the beginning of the solution, a relatively high CFL number will be accompanied by significant numerical smoothing. As the iteration count approaches the specified ending value, the CFL number will be gradually adjusted toward its final value as the amount of smoothing is steadily decreased. Details, as well as recommended values for the smoothing coefficients, are presented with the ACCELERATE keyword description (see p. 113).

For unstructured grids, only the CFL number variation and the zone are specified with the ACCELERATE keyword. No variation in smoothing is done; the same amount of smoothing is used for all iterations, as specified using the LAX DAMPING and TVD keywords. To reduce the amount of smoothing after the initial stages of the calculation, you must restart the solution with less dissipative smoothing values.

Keywords: ACCELERATE, LAX DAMPING, SMOOTHING, TEST 49 2, TVD

5 Boundary Conditions

The boundary conditions are perhaps the most important factor in influencing the accuracy of the flow computation. The manner in which the boundary conditions are imposed also influences the convergence properties of the solution. Wind-US uses a cell-vertex discretization, which results in solution points located on the boundaries of the zones which comprise the flow domain. During the computation, Wind-US computes the boundary values for the conservative variables, the species (if present), and turbulence variables (if present).

Proper specification of the flow boundary conditions is aided by a basic understanding of characteristic theory of the incoming and outgoing waves normal to the boundary. The boundary normal is considered positive when it points into the flow domain. The wave speeds (eigenvalues) have convective and acoustic components. A positive wave speed indicates a wave entering the flow domain, and so, a physical boundary condition must be specified and some auxiliary information must be supplied to impose the boundary condition. A negative eigenvalue indicates a wave leaving the flow domain, and so, a numerical boundary condition can be specified using flow data within the domain. The waves moving tangential to the boundary are neglected in the boundary condition treatment.

5.1 Explicit and Implicit Boundary Conditions

Wind-US imposes the boundary conditions explicitly after the interior solution has been computed for each zone after each iteration, by default. An exception is the zone interface boundaries, which are updated after each cycle. Another exception is the mass flow boundary condition, which is updated after a specified number of iterations (the default is five iterations) in order to reduce computational effort. Errors due to explicit boundary conditions are reduced through the use of the multi-stage or iterative time integration methods.

The use of implicit boundary conditions with an implicit solver is known to improve the stability of the method and lead to faster convergence at the expense of greater computational effort and complexity. Implicit boundary conditions are available in Wind-US in a limited manner, primarily for wall boundary conditions. The IMPLICIT BOUNDARY keyword is used to turn on implicit boundary conditions.

Keywords: IMPLICIT BOUNDARY

5.2 Boundary Condition Types

The Grid MANipulation (GMAN) program is used to associate boundary condition types with the solution points on the boundaries of the zones. These boundaries represent the geometry model, fluid boundaries, grid topological boundaries, and couplings between zones. The type of boundary conditions available in GMAN, in the order displayed by GMAN, include¹²:

- undefined
- reflection
- freestream
- viscous wall
- arbitrary inflow
- outflow

 $^{^{12}}$ In this User's Guide, GMAN boundary condition types are indicated by lower-case words in a fixed-pitch font, like this; Wind-US keywords are displayed in an upper-case fixed-pitch font, LIKE THIS.

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- inviscid wall
- self-closing
- singular axis
- coupled
- bleed
- pinwheel axis
- frozen
- chimera

It is possible to group some of these boundary condition types in a logical manner for detailed discussions, which are presented below. Some boundary conditions require further information, which is provided through keywords in the input data file, beyond being flagged in GMAN.

5.3 Wall Boundary Conditions

The inviscid wall, viscous wall, and bleed boundary condition types are all wall boundary conditions which simulate interaction of the flow with a real or imaginary solid surface.

5.3.1 Inviscid Wall

The inviscid wall boundary condition imposes flow tangency at the zone boundary (wall surface) while maintaining the same total velocity as the point adjacent to the boundary. One numerical boundary condition is imposed by computing the pressure at the boundary through an interpolation of interior pressures. A zero-order extrapolation is robust; however, the pressures may not be smoothly varying at the boundary. A first-order extrapolation works well for flows without discontinuity, and for flows in which the pressure does not vary greatly normal to the boundary. The extrapolations across a discontinuity may result in nonphysical pressures. One can use the viscous wall boundary condition along with the TURBULENCE INVISCID keyword in the input data file. This is useful if one wants to start a computation with the boundary as inviscid and then later turn on the viscous boundary conditions. In Wind-US, it is also possible to specify through TEST 138 that the normal pressure gradient at the wall be calculated rather than simply assuming it to be zero. Also, a first-order extrapolation which accounts for grid spacing can be used through the use of TEST 141.

5.3.2 Viscous Wall

The viscous wall boundary condition imposes a no-slip condition of the flow, a zero pressure gradient, and the appropriate heat transfer condition (adiabatic or constant temperature) at the zone boundary (wall surface). The no-slip condition can involve a non-zero velocity if the wall is moving. (See the MOVING WALL and ROLL keywords; however, for moving walls, the boundary condition type should be set to bleed.) To minimize transients at the start of a Wind-US calculation, the velocity at no-slip boundaries is actually reduced from its initial value to the no-slip condition over a number of iterations. The number of iterations may be specified using the WALL SLIP keyword.

The choice of the heat transfer condition is determined through the use of the WALL TEMPERATURE keyword in the input data file. The default is an adiabatic wall (zero temperature gradient). The temperature for the constant temperature condition is specified through the WALL TEMPERATURE keyword. The TTSPEC keyword is available for specifying a point-by-point distribution of surface temperatures.

Wall function boundary conditions may be used at viscous walls, using the White-Christoph law of the wall, through the WALL FUNCTION keyword. This feature is currently available for single-species flows only.

In Wind-US, it is also possible to specify through TEST 138 that the normal pressure gradient at the wall be calculated rather than simply assuming it to be zero. Also, a first-order extrapolation which accounts for grid spacing can be used through the use of TEST 141. Viscous flow is computed when the TURBULENCE keyword is used.

Keywords: MOVING WALL, TTSPEC, TURBULENCE, WALL FUNCTION, WALL SLIP, WALL TEMPERATURE

5.3.3 Bleed

The bleed boundary condition allows mass to flow through a porous, viscous wall. Bleed is mass flow out of the flow domain, while blowing is mass flow into the flow domain. Bleed and blowing systems are often an integral part of aeropropulsion configuration design, helping to control such flow phenomena as boundary layer growth and mixing. Wind-US's bleed / blowing boundary condition was designed to provide a means to model these systems with CFD. Specification of the bleed boundary condition in GMAN, which involves the identification of a particular bleed region number, triggers the calculation of the area of the bleed region specified, which is then stored in the grid file. Wind-US uses this area and the bleed or blowing conditions specified in the input data file to compute a normal velocity on the model surface. Bleed may be specified as a mass flow, or as a porous surface with a discharge coefficient and back pressure. Blowing may be specified through mass flow, plenum, or valve conditions.

Keywords: BLEED, BLOW

Moving Wall

The moving wall boundary condition enables a tangential velocity to be applied at no-slip walls in order to model rotating hubs or other components in the flow. The boundary solution points for the moving wall should be identified as bleed regions in GMAN. The translating or spinning motion of the wall is specified through the MOVING WALL keyword in the input data file. The ROLL keyword allows a rolling motion to be imposed on the grid.

Keywords: MOVING WALL, ROLL

5.4 Flow Interface Boundary Conditions

The freestream, arbitrary inflow, and outflow boundary condition types form a group involving the simulation of the interaction of the flow with other flow conditions at the domain boundaries.

5.4.1 Freestream

The freestream boundary condition is intended for use at freestream outer boundaries. This boundary condition uses one-dimensional characteristic theory to set boundary flowfield variables from freestream or flowfield conditions, based on the flow direction at the boundary.

At freestream boundaries with inflow, the HOLD keyword may be used to specify whether total conditions or characteristic values are to be held constant. Total pressure is held constant in both cases, and may result in initial Mach numbers being altered.

For some cases, this boundary condition may also be used at unconfined outflow boundaries, but the outflow boundary condition described in Section 5.4.3 is generally recommended instead. In particular, experience has shown that using the freestream condition at outflow boundaries with a shear layer exiting the computational domain may result in very slow convergence, and the solution may not be very accurate. There may also be the possibility of a mixed boundary, such as a supersonic outflow with a small subsonic region in the wall boundary layer. In this case, the pressure in the supersonic region is extrapolated to the boundary from upstream, but the pressure in the subsonic region is set from the freestream value. This may cause a small disturbance at the boundary, which can be corrected by specifying the boundary as an outflow boundary and imposing a constant pressure for the entire boundary.

Keywords: HOLD, EXTRAPOLATE

5.4.2 Arbitrary Inflow

The arbitrary inflow boundary condition allows conditions to be specified on regions of zonal boundaries where flow is entering the zone. Such a capability may be required to describe a thermally stratified nozzle input flow, or a jet emanating from a wall. The inflow profile may be specified in a number of different ways: as uniform flow, as a point-by-point (xyz) profile, or as uniform flow over a range of grid indices. The ARBITRARY INFLOW keyword is used in the input data file to indicate desired flow properties.

Keywords: ARBITRARY INFLOW, EXTRAPOLATE

5.4.3 Outflow

The outflow boundary condition may be used for internal flows at boundaries where subsonic flow is leaving the computational domain, such as at the exit plane of an inlet, diffuser, or auxiliary flow duct. It is also recommended for downstream outflow boundaries in external flow problems, especially if a shear layer is exiting the computational domain.

Characteristic theory indicates that only one physical condition is required to define the boundary condition. One may know one of the following at the outflow boundary: mass flow, exit pressure, or exit Mach number.

The mass flow (in lb_m/sec) may be specified at the outflow boundary (see the MASS FLOW keyword). The mass flow may be the actual or corrected value. One may alternatively specify the ratio of the desired mass flow to the mass flow through the inflow capture area specified in GMAN. During the solution, the mass flow boundary condition is applied every five iterations by default, which reduces computational costs. The integrated mass flow is compared with the desired value. If the PRESSURE option is used with the MASS FLOW keyword, a spatially-constant pressure is set at the outflow boundary, and modified as the solution proceeds until the desired mass flow is achieved. If the DIRECT option is used, the momentum, and thus the mass flow, is modified directly, and the pressure adjusts as the solution proceeds. For the PRESSURE option, Wind-US displays the ratio between the computational and desired mass flows and the modified pressure at each application of the boundary condition. If you experience difficulty in converging the mass flow, you should consider setting a constant back pressure at the duct exit.

A constant exit pressure may also be specified (see the DOWNSTREAM PRESSURE keyword) at the outflow boundary. This option results in a very reflective boundary condition, which may cause difficulties in convergence of the solution, especially for internal flows. One alternative is to allow the exit pressure to vary spatially according to the distribution of the solution points adjacent to the boundary. This option is selected through the VARIABLE option of the DOWNSTREAM PRESSURE keyword. The UNSTEADY option of the DOWNSTREAM PRESSURE keyword may be used to specify either a sinusoidal or user-defined pressure oscillation at an outflow boundary.

The mass-averaged Mach number may be specified at the outflow boundary using the DOWNSTREAM MACH keyword. This boundary condition is identical to the Chung-Cole compressor face boundary condition discussed below. It simulates the uniform Mach number characteristics that have been observed experimentally at compressor faces. This boundary condition also corresponds to a fairly uniform mass flux through the outflow boundary.

The compressor face models of Chung and Cole (1996) and of Mayer and Paynter (1994) are also available at outflow boundaries, through the COMPRESSOR FACE keyword. Both models are based on the observation that turbine engine conditions set the corrected mass flow, and that this corresponds directly to the average Mach number at the compressor face. These boundary conditions have been implemented mainly for the analysis of unsteady flow; however, they have also been shown to be robust for the establishment of steady-state, supercritical inlet flows.

The computational grid at the outflow boundary should be such that it is modeled with a single computational plane (constant i, j, or k) in a single zone. If two zones merge together near the exit, one should create a small exit zone to accommodate the outflow boundary condition.

Keywords: COMPRESSOR FACE, DOWNSTREAM MACH, DOWNSTREAM PRESSURE, MASS FLOW

5.5 Grid Topology Boundary Conditions

The reflection, self-closing, singular axis, and pinwheel axis types of boundary conditions form a group involving the simulation of the flow at topological surfaces in the grid.

5.5.1 Reflection

The reflection boundary condition simulates a plane of symmetry, and is the same as a solid, slip wall boundary condition. Therefore, within Wind-US, the inviscid wall boundary condition is actually applied. The reflection boundary condition type does provide a descriptive label for the boundary solution points, which may be of use to several auxiliary CFD codes which generate or use reflected grids.

5.5.2 Self-Closing

The self-closing boundary condition can be used at boundaries for which grid lines connect end-to-end (e.g., i_{max} connecting to i_1 as in an O-grid) in a point-match manner. The boundary condition simply averages the flow variables from both sides of the boundary and assigns the average to the two boundaries. This condition was formerly used for polar-type grids in duct flowfields. Note: For best results, one should consider using the coupled boundary condition instead of self-closing.

5.5.3 Singular Axis

The singular axis boundary condition is imposed at locations where an entire or part of a zonal boundary has collapsed to a line (not a point). Thus, along the other boundary direction, the grid points are coincident. The singular grid point is evaluated by taking the distance-weighted average of the solution of the adjacent grid points encircling the axis. This boundary condition is not to be used when the singularity line collapses to a point. For partially singular boundaries, the average is computed only over the singular portion of the boundary. Excessive use of this boundary condition is not recommended since flow conservation is not preserved. One should attempt to use non-singular grids whenever possible. TEST 118 allows the choice of which variables are averaged. TEST 150 can be used to explicitly set certain velocity components to zero when the singular axis occurs on symmetry planes. TEST 199 excludes the last grid point on the singular axis from being averaged.

5.5.4 Pinwheel

The pinwheel axis boundary condition is used when there exists multiple singularities at various locations on boundaries. This boundary condition does not zero any velocity components — it simply takes the average. TEST 118 allows the choice of which variables are averaged. TEST 165 allows input of the integer for the order of magnitude of the tolerance for singularity (the default is 10^{-8}).

5.6 Zonal Interface Boundary Conditions

The coupled and chimera boundary condition types involve the interactions between zones of the domain. For structured grids, periodic boundaries are available as a type of coupled boundary condition.

5.6.1 Coupled

The coupled boundary condition is imposed at regions where zones connect. Zone coupling is the name of the process by which Wind-US transfers flowfield information from one zone to another across conterminous computational grid boundaries. This process uses geometric interpolation factors stored in the grid file, which have been previously computed in the GMAN program. The zone interfaces occur at the boundaries of the zones, and so are imposed as boundary conditions at the end of each cycle of the computation.

The zone coupling algorithm that will be used depends on the explicit differencing operator specified by the RHS keyword. For most of the higher-order operators, the default zone coupling algorithm is a high-order method based on Roe's flux-difference splitting scheme, including the passing of gradients between zones. For structured grids the COUPLING keyword allows one to specify low-order Roe coupling, without passing gradients, or a coupling algorithm based on one-dimensional characteristic theory.

Zone boundaries should be treated the same as you would treat interior grid planes; there should not be any large changes in grid stretching or orthogonality at the boundary. In addition, zone boundaries should not be placed in regions of strong flowfield gradients, especially horizontally along jet shear layers or aligned with normal shocks. In other words, use your best engineering judgement in placing zone boundaries in your solution.

Periodic boundaries are currently supported only for structured grids. They are treated as normal coupled boundaries, with the connection data stored in the common grid (.cgd) file when setting boundary conditions with GMAN. See the PERIODIC "keyword" for details.

Keywords: COUPLING, PERIODIC

5.6.2 Chimera

The chimera boundary condition indicates that the zone boundary is in an overlap region.

5.7 Miscellaneous Boundary Conditions

The <u>undefined</u> and <u>frozen</u> boundary condition types are included in this section, mostly because they don't fit well in any other section, and are fairly self-explanatory.

5.7.1 Undefined

The default boundary condition type in GMAN is undefined. If not set to another boundary condition type, the boundary solution point is evaluated as an average of local solution points. In practice, one should specify the actual boundary condition type for all boundary solution points and avoid having undefined boundary solution points.

If all the points on a boundary surface have an undefined boundary condition, an error message is printed and the solution will abort. If only some points have an undefined boundary condition, a warning message is printed and the solution will continue.

5.7.2 Frozen

The frozen boundary condition simply signals a boundary solution point to retain its value as read in from the initial solution file.

6 Convergence Monitoring

Monitoring and properly assessing convergence levels during a Wind-US run are critical in obtaining meaningful, useful results. Wind-US allows you to track convergence by following residuals and/or integrated forces, moments, and mass flow. For engineering applications, the recommended convergence monitoring method is the tracking of integrated quantities of interest. For example, if you are modeling a wing/body geometry to determine drag, you should monitor integrated drag and set some reasonable bounds on drag oscillations as your convergence criterion.

6.1 Residuals

In steady-state (non-time-accurate) solutions, the residuals are the amounts by which the solution vector changes in a single iteration. Ideally, in approaching a steady state solution, the residuals should approach zero. However, in practice, complex geometric and flowfield features may limit the reduction in the residuals to about two orders of magnitude. Wind-US prints solution residuals to the list output file, in order to get a general idea of solution convergence.

For structured grids, Wind-US organizes the equations to be solved into logical "groups" that are solved together. For example, when a one- or two-equation turbulence model is used, the Navier-Stokes equations and the turbulence model equations are in two separate groups.

For each equation group in each zone, Wind-US prints the zone number, cycle number, location of the maximum residual (i, j, and k indices), equation number for which the maximum residual occurred, the value of the maximum residual, and the L2-norm of all the residuals for all the equations over all the points in that zone. The L2-norm of the residuals will give you an idea of the overall convergence of the solution. The location of the maximum residual may give you insight into problems with a particular solution. After solution "bombs" or when convergence is unacceptably slow, the location of the maximum residual is the first place you should look for potential problems with the solution.

The residuals are printed by default each iteration. The output interval may be changed, however, using the CYCLES and ITERATIONS keywords. Residuals may also be plotted by first using the auxiliary program *resplt* to generate a GENPLOT file, then using that file as input to the plot command in the CFPOST post-processing package.

By default, if the maximum residual for an equation group decreases by four orders of magnitude in a particular zone, a "converged" message is printed in the list output file, and iterations for that group in that zone are ended for the current cycle. Note that, if the residuals early in the calculation are large due to large initial solution transients, Wind-US may decide that a zone is converged when it may require significantly more iterations. In the input data file, with the CONVERGE keyword you can specify different convergence criteria, by setting the amount the residual must decrease, and by telling Wind-US to use the value of the residual rather than an order-of-magnitude reduction. You can also specify use of the L2-norm of the residual instead of the maximum residual using TEST 128.

Keywords: CONVERGE, CYCLES, ITERATIONS, TEST 128

6.2 Integrated Flowfield Quantities

As noted above, the optimal convergence checking criterion for a solution is an integrated quantity of interest for a particular study. For example, for an afterbody drag study to calculate pressure drag, the integrated drag is obviously the best quantity to track in monitoring convergence. This

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may be accomplished using Wind-US's flowfield integration capability. In the input data file, you may specify a number of computational surfaces for which Wind-US will integrate forces, moments, and/or mass flows. If you specify reference length, area, and moment center, Wind-US will output the forces and moments as coefficients; you may even specify output of lift and drag coefficients in place of x and y force coefficients.

You may specify zones and computational grid indices of subsets to be integrated, and you may request that different quantities be integrated on each subset. You may also control the frequency of output and whether the code outputs subset totals and/or zonal and grand totals. The auxiliary program *resplt* will produce GENPLOT plot files of output integration quantities versus solution cycle number, which may then be used as input to the **plot** command in the CFPOST post-processing package.

Keywords: LOADS

6.3 History Tracking of Flow Data

In time-accurate calculations, where a constant time step is being specified and is the same throughout the flow field, you may wish to track a particular flow variable as the solution advances in time. For example, you may be interested in monitoring the static pressure on a backward-facing step as vortices roll off the back of the step. You may request history tracking in the input data file. Specification must include zone numbers and ranges of grid points to track, and the frequency of the sampling. When you request history tracking, Wind-US creates a time history file containing the sampled data. The auxiliary program *thplt* must be run to extract data from the history file into GENPLOT plot files, which may then be used as input to the **plot** command in the CFPOST post-processing package.

Keywords: HISTORY

7 Files

A number of files are used by Wind-US in the course of a solution. The script file you run to submit Wind-US jobs will assign all the necessary files to their appropriate Fortran unit numbers, so you should not need to do any of that yourself. Each of the support files is described briefly below.

7.1 Input Data File (.dat)

Fortran unit number 5

The *input data file* is the primary control file for Wind-US and must be created for each case you want to run with the code. Input data and code options are entered through the use of descriptive keywords in this file. You should observe the following formatting rules in creating the input data file:

- 1. The first three lines of the file are reserved for geometry, flow condition, and arbitrary titles, respectively. Each of these titles may be up to 64 characters long. None of these first three lines may start with the (case-insensitive) word "Include".
- 2. Comment lines, beginning with a '/' character, may be placed anywhere in the file after the first three lines. Data file readability may be improved dramatically through the liberal use of comments for example, separating logical sections of the data file: numerical algorithm, force integrations, test options, etc.
- 3. Block data, such as that specified in arbitrary inflow or chemistry cases, must be contiguous—only keywords corresponding to the block may reside between the beginning and ending block indicators (e.g., CHEMISTRY and ENDCHEMISTRY).
- 4. Keywords may be entered in upper or lower case.
- 5. Abbreviations for keywords may be used, as long as they are unique. If they are not, you may not get the results you expect. For example, it's not a good idea to use single-letter abbreviations for keywords.

The following is an example of a simple input data file:

```
Geometric Title
Flowfield Condition Title
Optional Title
Freestream static 0.9 14.7 530. 4. 0.
Cycles 15
Inviscid
End
```

7.2 Grid File (.cqd)

Fortran unit number 11

The computational grid used by Wind-US for a particular case is stored in the *grid file*. In this file are stored the (x, y, z) coordinates of all computational grid points, zone coupling interpolation factors, and grid reference and scaling data.

This file was originally referred to as the *common* grid file, so named because the file was formatted according to Boeing's Common File Format (CFF). Wind-US also supports grid files in

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CGNS (CFD General Notation System) format.¹³ Both common files and CGNS files are binary, and portable to virtually every hardware platform, except Cray, with no need for explicit data conversions.

Grid files for Wind-US may be created by several mesh-generation codes in either common file or CGNS format. If necessary, the *cfcnvt* utility may be used to convert a variety of other formats, including PLOT3D xyz format, to common file format.

Zone coupling, reference, and scaling data are added to common grid files using the GMAN program. Common grid files are also used in the CFPOST post-processing package. Neither GMAN nor CFPOST currently support CGNS files, however.

Keywords: CGNSBASE

7.3 Flow File (.cfl)

Fortran unit number 20

The *flow file* contains the computed flow field. For Navier-Stokes and Euler solutions, the file contains density, momentum, and energy data, and, for viscous solutions, turbulence data. The flow file also records the current solution cycle number to allow the file to be used for solution restarts.

Like the grid file, the flow file was originally referred to as the *common* flow file, but may now be written in either common file or CGNS format. Several graphical post-processing programs are able to read both common files and CGNS files.

The CFPOST post-processing package may be used with common flow files to produce other files for post-processing and/or to create flowfield plots directly. CFPOST does not currently support CGNS files, however.

Keywords: CGNSBASE

7.4 Boundary Data File (.tda)

Fortran unit number 14

Wind-US's boundary data file is used during solution restarts, and results in smoother restarts, especially with higher-order boundary coupling. The file acts as a buffer for the transfer of zone coupling information and a holding bin for data needed throughout a Wind-US run but not stored in the grid or solution file.

7.5 Time History File (.cth)

Fortran unit number 19

The time history file is a common file which stores data resulting from the use of Wind-US's history tracking capability. The file contains a lookup table corresponding to the range of computational indices tracked during the run, time stamp data, and flowfield data for each time stamp. Upon completion of a time history run, the auxiliary program thplt may be used to view the contents of the time history file.

Keywords: HISTORY

¹³Detailed information on the CGNS standard may be found at the CGNS web site, at http://cgns.sourceforge.net/.

7.6 List Output File (.lis)

Fortran unit number 6

The purpose of Wind-US's list output file is to echo the input from the input data file, track convergence and integration results, record CPU/job statistics, and log code error messages. The auxiliary program resplt may be used to extract convergence and integration data from the list output file and create a GENPLOT-style data file, which may be plotted with the plot command in the CFPOST post-processing package. For parallel-processing runs, this file also contains messages relating to the PVM system and to the allocation and operation of slave processors. Convergence data is written to this file for each of the equation sets solved during the run, including the Euler, Navier-Stokes, and non-algebraic turbulence model sets.

7.7 Time Data File (.cft)

Fortran unit number 22

The time data file may be used for storing the computed flow field at the extra time levels required for second-order time differencing and/or Newton iteration. Currently by default, the extra time levels for second-order time differencing are stored in the .cfl file, and those for Newton iteration are stored in the .cft file and linked to the .cfl file. Keywords in the TEMPORAL keyword block may be used to specify where the extra time levels should be stored.

Keywords: IMPLICIT ORDER, NEWTON, TEMPORAL

7.8 Edge Data File (.cge)

Fortran unit number 21

The edge data file is used with unstructured grids to store grid-related information about each cell that's required during the solution procedure. The file is created during the initial run, and by default is saved for use in subsequent restart runs, but will be recreated if it's missing. When the file is opened, if necessary it is automatically split into multiple files to keep the size of each file below two gigabytes, with the main file transparently linked to the separate files.

Keywords: DEBUG 8, DEBUG 9

7.9 Wind-US Stop File (NDSTOP)

Fortran unit number 1

The stop file is used to stop Wind-US execution cleanly in the middle of a CFD solution. Although a solution may be stopped simply by killing the Wind-US process from the system queue, doing so will not ensure a clean update of all zonal flowfield data to the flowfield file. The stop file — named NDSTOP — provides a means to cleanly shut down a running solution, completing the current cycle or zone, performing zone coupling, updating the flowfield file, and removing all symbolic links to Fortran file unit numbers.

To stop the code, you must place one of two words in the stop file in the directory in which the Wind-US job is running. The word STOP in the stop file signals Wind-US to complete the current cycle (at the end of the last zone) and exit. In single-processing mode, you may also use the word

STOPZONE in the stop file to stop Wind-US after completing the zone currently in memory.¹⁴ You may then restart the code in your next run, starting at the next zone or back at zone one. Regardless of the existence of this file, Wind-US will stop if the requested number of cycles has been computed, or if your solution has converged.

During the clean-up procedure at the end of the job, the NDSTOP file is automatically removed.

Note that the directory in which the Wind-US job is running, where the *NDSTOP* file must be, is not necessarily the one you were in when the *wind* command was issued. If you don't use the *-runinplace* option to the *wind* script (see Section 8.1), the job will be run in a remote directory. The root name of the remote run directory may be specified using the *-runroot* option, or in response to a command line prompt. The default for the "root name" of the remote directory is /tmp. The full name of the remote directory will be rootname/userid/basename.scr, where rootname is the root name described above, userid is your userid, and basename is the base name of your .dat file.

On a Unix system, you might submit a simple 'at' job for a later time as follows:

```
at 0530 monday
echo STOP > NDSTOP
^D
```

At 5:30 AM on the next Monday, the system would create the *NDSTOP* file with the word 'STOP' in it. Note that this will not stop the run exactly at 5:30 AM; Wind-US must still complete the current cycle, which may take an hour or more for large cases.

Keywords: RESTART

7.10 Temperature and Transition Specification Files

Fortran unit number 45

Wind-US can read temperature and transition files, specifying the temperature or transition to turbulence on any boundary surface, that were created with the older *tmptrn* utility. This option is included for backward compatibility with WIND 2.0, and is only needed for an initial (i.e., non-restart) run. The *wind* script copies the files to the run directory, and Wind-US opens the files directly (unit 45) using the file names specified with the TTSPEC keyword.

New applications should use the latest tmptrn utility to write the temperature/transition data into the common flow (.cfl) file directly.

Keywords: TTSPEC

7.11 Chemistry Files (.chm)

Fortran unit number 26

The generalized chemistry files contain all the information Wind-US requires to compute a general chemistry mixture. A chemistry file has a header line, followed by three sections containing data defining the thermodynamic properties, reaction rates, and transport properties.

¹⁴In multi-processing mode, STOPZONE does the same thing as STOP.

7.11.1 Header

ispec

The first line in the file must contain a single value, ispec, specifying the type of reaction and the format for the finite-rate data. The format for the line is 15 :

where	
\underline{ispec}	Reaction and Format
100, 110, 115	Forward and backward elementary reactions, with two reactants and two products. Backward rate computed using an equilibrium constant. (Format 1)
130, 135, 136, 137	Forward and backward elementary reactions, with two reactants and two products. Forward and backward rates may be specified separately, or one may be computed using an equilibrium constant. For $ispec=137$, in addition to elementary reactions with two reactants and two products, "general" and "global" exchange reactions with up to three reactants and three products may also be used. (Format 2)
120, 121	Forward reactions only. Intended for detonation or rapid combustion problems. (Format 3)

It should be noted that *ispec* values of 1, 3, and 4 were used in earlier versions of Wind-US. Values of 1 and 4 are now equivalent to ispec = 100, and 3 is now equivalent to ispec = 130.

7.11.2 Thermodynamic Properties

ISPEC

This section of the chemistry data file contains the information necessary to compute the thermodynamic properties for each species. The general format for this section is:

```
THERMODYNAMIC COEFFICIENTS

Title, line 1

Title, line 2

ns NS

Curve type

Information defining species 1

...

Information defining species ns
```

where *ns* is the number of species, and *Curve type* specifies the type of curve fits used to define the thermodynamic properties. The section title THERMODYNAMIC COEFFICIENTS must be the second line in the file, immediately after the line defining *ispec*. The *Curve type* line may be omitted (not left blank), or specified as one of SPARKCRV, WINDNASA, and NASA3287. If the line is omitted, the SPARKCRV curve type is assumed.

The information defining each species, and the specific format used, depends on the curve type, as described below.

¹⁵The value *ispec* is read using a list-directed read statement. The ISPEC label following the value is thus optional, but present in the standard chemistry files supplied with Wind-US. The labels following the values *ns*, *nreq*, *ndeq*, and *tfrmin*, described in Section 7.11.2 and Section 7.11.3, are also optional.

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This is the original file format used for Wind-US chemistry files. The specific heat at constant pressure C_p for each species is defined by a series of fourth-order polynomials, each valid within a defined temperature range. I.e.,

$$\frac{C_p}{R} = a_1 + a_2T + a_3T^2 + a_4T^3 + a_5T^4$$

where R is the universal gas constant, and T is in K.

Given the above, polynomials may be derived for the enthalpy, entropy, and Gibb's free energy.

$$\frac{H}{RT} = a_1 + a_2 \frac{T}{2} + a_3 \frac{T^2}{3} + a_4 \frac{T^3}{4} + a_5 \frac{T^4}{5} + \frac{b_1}{T}$$

$$\frac{S}{R} = a_1 \ln T + a_2 T + a_3 \frac{T^2}{2} + a_4 \frac{T^3}{3} + a_5 \frac{T^4}{4} + b_2$$

$$\frac{G}{RT} = a_1 (1 - \ln T) - a_2 \frac{T}{2} - a_3 \frac{T^2}{6} - a_4 \frac{T^3}{12} - a_5 \frac{T^4}{20} + \frac{b_1}{T} - b_2$$

For the SPARKCRV curve type, the enthalpy reference state is at T=0 K. I.e., the curve fit for H/RT actually computes

$$\frac{H}{RT} = \frac{\Delta_f H_{(0)} + \Delta H_{(0 \to T)}}{RT}$$

where $\Delta_f H_{(0)}$ is the heat of formation at 0 K, and $\Delta H_{(0\to T)}$ is the change in enthalpy between 0 K and T.

For each species, the information defining the species and for computing C_p/R , etc., is stored in the .chm file as described in the following table. Note that records 2–3 are repeated for each temperature range.

Contents of Record	Columns	Format
Record 1		
• Name of species	1-8	a8
• Number of curves (i.e., temperature ranges) defining thermodynamic properties	16–20	i5
• Name, and number per molecule, of constituent elements	25 – 44	4(a2,f3.0)
• Low-temperature (i.e., constant) value for C_p/R The default is the value computed from the polynomial curve fit, evaluated at the beginning temperature for the first curve.	51–60	f10.1
Molecular weight	66 - 75	f10.3
Record 2		
• Minimum and maximum temperature for curve	1 - 30	2e15.5
• Coefficients a_1 – a_3 in equations for thermodynamic properties	31 - 75	3e15.5
Record 3		
• Coefficients a_4 , a_5 , b_1 , and b_2 in equations for thermodynamic properties	1–60	4e15.5

Example

THERMODYNAMIC COEFFICIENTS

FROM NASA-RFL-TR-70-3, NASA-CR-111989, MAC LIB FISH NO. N71-38747

5 NS				
SPARKCRV				
02	3 0 2.	0. 0. 0.	3.5	32.000
300.000	1000.000	0.37190E+01	-0.25170E-02	0.85840E-05
-0.83000E-08	0.27080E-11	-0.104419E+04	0.00000E+00	
1000.000	6000.000	0.33160E+01	0.11510E-02	-0.37260E-06
0.61860E-10	-0.36660E-14	-0.104419E+04	0.00000E+00	
6000.000	15000.000	0.37210E+01	0.42540E-03	-0.28350E-07
0.60500E-12	-0.51860E-17	-0.104419E+04	0.00000E+00	
NO	3 0 1.N	1. 0. 0.	3.5	30.008
300.000	1000.000	0.41470E+01	-0.41200E-02	0.96920E-05
-0.78630E-08	0.22310E-11	0.979001E+04	0.00000E+00	
1000.000	6000.000	0.32210E+01	0.12210E-02	-0.42970E-06
0.65590E-10	-0.34510E-14	0.979001E+04	0.00000E+00	
6000.000	15000.000	0.38450E+01	0.25210E-03	-0.26580E-07
0.21620E-11	-0.63810E-16	0.979001E+04	0.00000E+00	
\dots etc.				

WINDNASA

This is the same as the SPARKCRV curve fits described above, except for the addition of a single final record for each species that specifies the heat of formation at 0 K. The same equations are used for C_p/R , etc. However, with the WINDNASA curve fits, the enthalpy (and Gibb's free energy, since G/RT = H/RT - S/R) use an enthalpy reference state of 298.15 K instead of 0 K. I.e., the curve fit for enthalpy actually computes

$$\frac{H'}{RT} = \frac{\Delta_f H_{(298)} + \Delta H_{(298 \rightarrow T)}}{RT}$$

where $\Delta_f H_{(298)}$ is the heat of formation at 298.15 K, and $\Delta H_{(298 \to T)}$ is the change in enthalpy between 298.15 K and T.

In Wind-US, when ADJUST is specified in the CHEMISTRY keyword block, the values of enthalpy and Gibb's free energy returned by the WINDNASA curve fits are shifted to change the reference state to 0 K. I.e.,

$$\begin{split} \frac{H}{RT} &= \frac{H'}{RT} + \frac{H_{(shift)}}{RT} \\ &= \frac{\Delta_f H_{(298)} + \Delta H_{(298 \to T)}}{RT} + \frac{\Delta_f H_{(0)} - \Delta_f H_{(298)} + \Delta H_{(0 \to 298)}}{RT} \\ &= \frac{\Delta_f H_{(0)} + \Delta H_{(0 \to T)}}{RT} \end{split}$$

When ADJUST is specified, $\Delta_f H_{(0)}$, the heat of formation at 0 K in the above equation, is read from the .chm file. If ADJUST is not specified, the enthalpy reference state is left as 298.15 K, and the line in the .chm file containing the heat of formation at 0 K, if present, is ignored.

For each species, the information defining the species and for computing C_p/R , etc., is stored in the *.chm* file as described in the following table. Note that records 2–3 are repeated for each temperature range. Except for the last record specifying the heat of formation at 0 K, this is the same as for the SPARKCRV format.

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Contents of Record	Columns	Format
Record 1		
• Name of species	1-8	a8
• Number of curves (i.e., temperature ranges) defining thermodynamic properties	16-20	i5
• Name, and number per molecule, of constituent elements	25 – 44	4(a2,f3.0)
• Low-temperature (i.e., constant) value for C_p/R The default is the value computed from the polynomial curve fit, evaluated at the beginning temperature for the first curve.	51–60	f10.1
• Molecular weight	66 - 75	f10.3
Record 2		
• Minimum and maximum temperature for curve	1 - 30	2e15.5
• Coefficients a_1 – a_3 in equations for thermodynamic properties	31 - 75	3e15.5
Record 3		
• Coefficients a_4 , a_5 , b_1 , and b_2 in equations for thermodynamic properties	1–60	4e15.5
Record 4 (only required if ADJUST is used)		
• Identifier string starting with the word "Heat"	1 - 31	a31
• Heat of formation at 0 K, in J/mole	32–46	f15.3

Example

THERMODYNAMIC COEFFICIENTS

CURVE FITS FROM NASA LEWIS CET86 HIGH TEMPERATURE THERMO DATA BASE

```
7
          NS
WINDNASA
02
                        0 2.
                                0.
                                     0.
                                          0.
                                                         3.5
                                                                     31.998
                      5000.000
        300.000
                               3.1162949E+00
                                              1.5886094E-03 -6.7904360E-07
  1.4714899E-10 -1.1729212E-14 -9.9401794E+02
                                               6.4600671E+00
       5000.000
                     15000.000 2.5782323E+00
                                              8.5796324E-04 -7.6397647E-08
  1.3412616E-12 3.2564804E-17
                                1.1504711E+03 1.1400551E+01
Heat of Formation at 0 deg K
                                0.0
Н
                                     0.
                                                                      1.008
                   2
                        H 1.
                                0.
                                          0.
                                                         2.5
        300.000
                      5000.000 2.5000000E+00 0.0000000E+00
                                                             0.000000E+00
 0.0000000E+00 0.0000000E+00 2.5474038E+04 -4.5991986E-01
       5000.000
                     15000.000 2.5000000E+00 0.0000000E+00
                                                             0.000000E+00
 0.0000000E+00 0.0000000E+00 2.5474038E+04 -4.5991986E-01
Heat of Formation at 0 deg K
                                216024.1
\dots etc.
```

NASA3287

This format for the curve fits defining the thermodynamic properties is derived from the one defined in NASA TP 3287 (McBride, Gordon, and Reno, 2001). A series of curves is again used to define the specific heat at constant pressure, enthalpy, entropy, and Gibb's free energy, with each

curve valid within a defined temperature range. I.e.,

$$\begin{split} \frac{C_p}{R} &= a_1 T^{-2} + a_2 T^{-1} + a_3 + a_4 T + a_5 T^2 + a_6 T^3 + a_7 T^4 + a_8 T^5 \\ \frac{H}{RT} &= -a_1 T^{-2} + a_2 T^{-1} \ln T + a_3 + a_4 \frac{T}{2} + a_5 \frac{T^2}{3} + a_6 \frac{T^3}{4} + a_7 \frac{T^4}{5} + a_8 \frac{T^5}{6} + \frac{b_1}{T} \\ \frac{S}{R} &= -a_1 \frac{T^{-2}}{2} - a_2 T^{-1} + a_3 \ln T + a_4 T + a_5 \frac{T^2}{2} + a_6 \frac{T^3}{3} + a_7 \frac{T^4}{4} + a_8 \frac{T^5}{5} + b_2 \\ \frac{G}{RT} &= -a_1 \frac{T^{-2}}{2} - a_2 T^{-1} (1 + \ln T) + a_3 (1 + \ln T) - a_4 T - a_5 \frac{T^2}{2} - a_6 \frac{T^3}{12} - a_7 \frac{T^4}{20} - a_8 \frac{T^5}{30} + \frac{b_1}{T} - b_2 \end{split}$$

where R is the universal gas constant, and T is in K.

Like the WINDNASA curve fits, the enthalpy reference state for the NASA3287 curve fits is 298.15 K. When ADJUST is specified in the CHEMISTRY keyword block, the values of enthalpy and Gibb's free energy are shifted to change the reference state to 0 K, as described above for the WINDNASA curve fits.

For each species, the information defining the species and for computing C_p/R , etc., is stored in the *.chm* file as described in the following table. Note that records 3–5 are repeated for each temperature range.

Contents of Record	Columns	Format
Record 1		
• Name of species	1-18	a18
• Comments (not used in Wind-US)	19 - 76	a58
Record 2		
• Number of curves (i.e., temperature ranges) defining thermodynamic properties	1–2	i2
• Identification code (not used in Wind-US)	4-9	a6
• Name, and number per molecule, of constituent elements	11 – 50	5(a2,f6.2)
• Flag indicating standard state (not used in Wind-US)	52	i1
• Molecular weight	53 – 65	f13.5
• Heat of formation at 298.15 K, in J/mole	66-80	f15.3
Record 3		
• Minimum and maximum temperature for curve	2-21	2f10.3
• Number of coefficients a_n in equation for C_p/R	23	i1
• Exponents of T in equation for C_p/R	24 – 63	8f5.1
\bullet Enthalpy difference from 298.15 K to 0 K, in J/mole	66-80	f15.3
Record 4		
• Coefficients a_1 – a_5 in equations for thermodynamic properties	1-80	5d16.8
Record 5		
• Coefficients a_6 – a_8 in equations for thermodynamic properties	1-48	3d16.8
\bullet Coefficients b_1 and b_2 in equations for thermodynamic properties	49 – 80	2d16.8
Record 6 (only required if ADJUST is used)		
• Identifier string starting with the word "Heat"	1 - 31	a31
• Heat of formation at 0 K, in J/mole	32 – 46	f15.3

THERMODYNAMIC COEFFICIENTS

Example

```
CURVE FIT JANAF 6000K DATA EXTRAP. TO 15000K & NASA LEWIS DATA RE-FITTED
    5
          NS
NASA3287
C02
                 Props & Hf298: TPIS v2,pt1,1991,p27.
 3 1 7/88 C
             1.000
                     2.00
                             0.00
                                     0.00
                                             0.00 0
                                                                   -393510.000
                                                       44.00980
    200.000 1000.000 7 -2.0 -1.0 0.0
                                      1.0
                                           2.0 3.0 4.0 0.0
                                                                      9365.469
 4.94378364D+04 -6.26429208D+02 5.30181336D+00 2.50360057D-03 -2.12470010D-07
 -7.69148680D-10 2.84997991D-13 0.0000000D+00 -4.52818986D+04 -7.04876965D+00
  1000.000 6000.000 7 -2.0 -1.0 0.0 1.0 2.0 3.0 4.0 0.0
                                                                      9365.469
 1.17696943D+05 -1.78880147D+03 8.29154353D+00 -9.22477831D-05 4.86963541D-09
 -1.89206384D-12 6.33067509D-16 0.0000000D+00 -3.90834501D+04 -2.65268192D+01
  6000.000 20000.000 7 -2.0 -1.0 0.0 1.0 2.0 3.0 4.0 0.0
 -1.54440594D+09 1.01683595D+06 -2.56137666D+02 3.36936340D-02 -2.18115756D-06
 6.99132366D-11 -8.84221052D-16 0.0000000D+00 -8.04312703D+06 2.25415288D+03
Heat of Formation at 0 deg K
                               -393149.56
H20
                 CODATA, 1989. JRNBS v92, 1987, p35. TRC tuv-25, 10/88.
 2 1 8/89 H
             2.000
                     1.00
                             0.00
                                     0.00
                                             0.00 0
                                                       18.01528
                                                                   -241826.000
    200.000 1000.000 7 -2.0 -1.0 0.0 1.0 2.0 3.0 4.0 0.0
                                                                      9904.092
 -3.94795999D+04 5.75572977D+02 9.31783351D-01 7.22271091D-03 -7.34255448D-06
 4.95504134D-09 -1.33693261D-12 0.0000000D+00 -3.30397425D+04 1.72420539D+01
  1000.000 6000.000 7 -2.0 -1.0 0.0 1.0 2.0 3.0 4.0 0.0
                                                                      9904.092
 1.03497224D+06 -2.41269895D+03 4.64611114D+00 2.29199814D-03 -6.83683007D-07
 9.42646842D-11 -4.82238028D-15 0.0000000D+00 -1.38428625D+04 -7.97815119D+00
```

7.11.3 Reaction Rates

 \dots etc.

This section of the .chm file contains the reaction rate information. The first five lines are:

-238918.95

```
FINITE RATE COEFFICIENTS Title, line\ 1 Title, line\ 2 nreq\ ndeq NREQ, NDEQ tfrmin TFRMIN
```

Heat of Formation at 0 deg K

where *nreq* is the total number of reactions (dissociation + exchange + ionization), *ndeq* is the number of dissociation reactions (i.e., have a third body), and *tfrmin* is the temperature in K below which no reactions occur. The section title FINITE RATE COEFFICIENTS must immediately follow the last data line in the "Thermodynamic Properties" section, with no blank lines in between.

There are three possible formats for the remainder of this section, depending on the value of ispec.

Format 1; ispec = 100, 110, 115

This option may be used for any chemically reacting flow with dissociation and exchange reactions of the form

$$S_1 \rightleftarrows S_3 + S_4$$

$$S_1 + S_2 \rightleftarrows S_3 + S_4$$

where S_1 , etc., represent chemical species.

Rate Equations. For dissociation reactions, the forward and backward reaction rate equations are of the form

$$r_f = k_f[S_1]$$
$$r_b = k_b[S_3][S_4]$$

where $[S_1]$ represents the concentration of species S_1 , etc.

For exchange reactions, the forward and backward reaction rate equations are of the form

$$r_f = k_f[S_1][S_2]$$
$$r_b = k_b[S_3][S_4]$$

Rate Coefficients. The forward and backward reaction rate coefficients k_f and k_b are computed using the equations

$$k_f = CT^S e^{-D/(K_B T)}$$
$$k_b = k_f/K$$

where T is the temperature in K, the ratio D/K_B is in K, and K_B is the Boltzmann constant. The coefficient C has units $(\text{cm}^3/\text{g-mol})(\text{sec-K}^S)^{-1}$.

For ispec = 100 or 110, the equilibrium constant K is computed using

$$K = \exp(a_1 + a_2 Z + a_3 Z^2 + a_4 Z^3 + a_5 Z^4)$$
$$Z = 10,000/T$$

For ispec = 115 the equilibrium constant K is computed from the change in Gibb's free energy for the reaction.

$$K = e^{-\Delta G^{\circ}/RT}$$

where ΔG° is the change in Gibb's free energy, R is the gas constant, and T is the temperature. The Gibb's free energy for each species is determined from the information specified in Section 7.11.2.

Third Body Reaction Rates. For the dissociation reactions the specification of the third body reaction rates depends on the *ispec* value used.

ispec Third Body Reaction Rate	ispec	Third	Body	Reaction	Rate
--------------------------------	-------	-------	------	----------	------

- Variable third body reaction rates are used. The .chm file contains the names of the third body reactants and the corresponding coefficients C used in the formula for k_f . Note that values must be specified for each of the ns species.
- 110, 115 Variable or average third body reaction rates are used. The .chm file contains the average third body coefficient C, and, for variable reaction rates, the names of the third body reactants and the corresponding third body efficiencies.

File Format. The information needed for each reaction is stored in the chemistry data (.chm) file, with the dissociation reactions listed first. I.e., reaction data is specified in the following order:

Information defining dissociation reaction 1

. . .

Information defining dissociation reaction ndeq Information defining exchange reaction 1

. . .

Information defining exchange reaction nreq - ndeq

where nreq is the total number of reactions, and ndeq is the number of dissociation reactions.

The data to be specified in the .chm file is described in detail in the following table.

Contents of Record	Columns	Format
Record 1 (for all ispec values)		
• Name of first reactant	1-5	a 5
• Name of second reactant (blank for dissociation)	9-13	a 5
• Name of first product	17 - 21	a 5
• Name of second product	25 – 29	a 5
• Temperature exponent S in equation for k_f	33 – 44	e12.4
• D/K_B in equation for k_f	45 - 56	e12.4
• Coefficient C in equation for k_f (only for $ispec = 115$). For dissociation reactions, this is the average third body coefficient.	57–68	e12.4
Record 2 (for all ispec values except 115)		
• Coefficients a_1 – a_5 in equation for K	1-60	5e12.4
$Records \ 2+ \ (for \ ispec = \ 110)$		

• For dissociation reactions with variable third body reaction rates, these records specify the third body reactants and the corresponding third body efficiencies. To use an average third body reaction rate with ispec=110, simply omit these lines.

Only the non-unity efficiencies have to be specified. The first record in this group is

```
THIRD [BODY] [EFFICIENCY] nnot1
```

where *nnot1* is the number of third body reactants with a non-unity efficiency. (The brackets are used to indicate that the words BODY and EFFICIENCY are optional.) The names of the third body reactants and the corresponding efficiencies are specified in the next *nnot1* records, in free format, as follows:

```
species(1) efficiency(1)
...
species(nnot1) efficiency(nnot1)
```

Continued on next page

(Continued from previous page)

Contents of Record Columns Format

Records 2+ (for ispec = 115)

 \bullet For dissociation reactions, these records specify the third body reactants and the corresponding third body efficiencies. The format is the same as described above for ispec=110, with one exception. The first record

THIRD [BODY] [EFFICIENCY] nnot1

must be specified even when an average third body reaction rate is to be used. To use an average third body reaction rate with ispec = 115, specify a zero value (not blank) for nnot1, and omit the lines listing the species and the corresponding efficiencies.

Records 3+ (for ispec = 100)

• Name of third body reactant (only for dissociation reactions)	17 - 24	a8
• Coefficient C in equation for k_f . For the dissociation reactions,	25 - 36	e12.4
this is the coefficient for the specified third body reactant.		

Record 3 (for ispec = 110)

• Coefficient C in equation for k_f . For the dissociation reactions, 25–36 e12.4 this is the average third body coefficient.

With Format 1, the data is read in subroutine frtin1, and the rates are computed in rates (for ispec = 100), rates a (ispec = 110), or rates and (ispec = 115).

Example

FINITE RATE COEFFICIENTS FROM AIAA 88-0513

5,3 2000.	NREQ,NDEQ TFRMIN				
02	0	0	-1.0	5.9500E+04	
1.335	-4.127	-0.616	0.093	-0.005	5
	02	2.75E+19			
	NO	2.75E+19			
	0	8.25E+19			
	N	8.25E+19			
	N2	2.75E+19			
N2	N	N	-1.6	1.1320E+05	
3.898	-12.611	0.683	-0.118	0.006	5
	02	3.70E+21			
	NO	3.70E+21			
	0	1.11E+22			
	N	1.11E+22			
	N2	3.70E+21			
NO	N	0	-0.5	7.5500E+04	
1.549	-7.784	0.228	-0.043	0.002	5

Format 2; ispec = 130, 135, 136, 137

This format is the most flexible one in Wind-US, with a variety of options for specifying the necessary information. "Elementary" dissociation and exchange reactions may be specified using $ispec=130,\ 135,\ and\ 136,\ the\ same$ as with Format 1. In addition, for ispec=1037, "general" and "global" exchange reactions may be specified with up to three reactants and products.

Rate Equations. General exchange reactions are of the form

$$c_1S_1 + c_2S_2 + c_3S_3 \rightleftharpoons c_4S_4 + c_5S_5 + c_6S_6$$

with forward and backward reaction rate equations of the form

$$r_f = k_f [S_1]^a [S_2]^b [S_3]^c$$
$$r_b = k_b [S_4]^d [S_5]^e [S_6]^f$$

Global exchange reactions are forward only, of the form

$$c_1S_1 + c_2S_2 + c_3S_3 \rightarrow c_4S_4 + c_5S_5 + c_6S_6$$

with corresponding reaction rate equations

$$r_f = k_f [S_1]^a [S_2]^b [S_3]^c$$
$$r_b = 0$$

Rate Coefficients. For all the *ispec* values, the forward and backward reaction rate coefficients k_f and k_b are computed using the equations

$$k_f = C_f T^{S_f} e^{-D_f/(K_B T)}$$

$$k_b = C_b T^{S_b} e^{-D_b/(K_B T)}$$

where T is the temperature in K, the ratios D_f/K_B and D_b/K_B are in K, and K_B is the Boltzmann constant. The reaction rate coefficients for the j'th reaction, $(C_f)_j$ and $(C_b)_j$, have units

$$(C_f)_j$$
: $\left(\frac{\mathrm{cm}^3}{\mathrm{g\text{-}mol}}\right)^{O_j-1} \frac{1}{\mathrm{sec\text{-}K}^{S_f}}, \qquad (C_b)_j$: $\left(\frac{\mathrm{cm}^3}{\mathrm{g\text{-}mol}}\right)^{O_j-1+\nu_j} \frac{1}{\mathrm{sec\text{-}K}^{S_b}}$

where O_j is the order of the reaction (i.e., the total number of moles of reactants), and ν_j is the number of moles of products minus the number of moles of reactants.

Note that unlike Format 1, with Format 2 the backward reaction rate coefficient k_b may be specified independently of the forward reaction rate coefficient. However, if the specified input value of either the forward or backward coefficient C_f or C_b is zero, the corresponding reaction rate coefficient k_f or k_b is computed from the non-zero reaction rate coefficient and an equilibrium constant that's computed using Gibb's function, just as in Format 1 with ispec = 115. E.g., if the specified $C_b = 0$, the backward reaction rate coefficient is computed as

$$k_b = k_b/K$$

where the equilibrium constant K is

$$K = e^{-\Delta G^{\circ}/RT}$$

Eddy Dissipation Concept. For ispec = 137, the eddy dissipation concept may be used, specified using the EDC keyword in the CHEMISTRY keyword block. Note that although three reactants and products are allowed with ispec = 137, the eddy dissipation concept should only be used with two reactants and forward reactions only, and is only applicable to non-premixed combustion. It's intended for rapid combustion reactions with the fuel as species S_1 and the oxidizer as species S_2 .

With the eddy dissipation concept, a second forward reaction rate is computed in addition to the one described above. The two reaction rates are compared, and the lower rate is used.

The reaction rate equation using the eddy dissipation concept is of the form

$$r = C_{edc} \frac{\rho}{M_1} c_{\mu} \omega \min\left([S_1], [S_2] \frac{c_1 M_1}{c_2 M_2} \right)$$

where C_{edc} is the eddy dissipation coefficient, specified using the EDC keyword in the CHEMISTRY keyword block; ρ is the fluid density; M_1 and M_2 are the molecular weights of species 1 (fuel) and species 2 (oxidizer); c_{μ} is a constant equal to 0.09; ω is the specific dissipation rate from the SST turbulence model; $[S_1]$ and $[S_2]$ are the concentrations of species 1 (fuel) and species 2 (oxidizer); and c_1 and c_2 are the number of molecules in the reaction for species 1 (fuel) and species 2 (oxidizer).

Note that the eddy dissipation concept may currently be used only with the SST turbulence model.

Damkohler Rate Limiter. Also for *ispec* = 137, the Damkohler rate limiter may be used, specified using the DAMKOHLER keyword in the CHEMISTRY keyword block.

With the Damkohler limiter, both forward and backward reaction rates are limited such that the ratio $Da = \tau_{FD}/\tau_{CH}$ is less than some user-specified maximum Da_{max} . The fluid dynamic time scale τ_{FD} is taken as the local time step Δt . The chemical reaction time scale τ_{CH} is the value that would allow the concentration of the specie with the smallest negative defect (i.e., the one that will first reach zero concentration) to drop by a ratio of 1/e.

Third Body Reaction Rates. Like in Format 1, for the dissociation reactions the specification of the third body reaction rates depends on the *ispec* value used.

\underline{ispec}	Third Body Reaction Rate
130, 136	Average third body reaction rates are used. The .chm file contains the average
	third body coefficients C_f or C_b .
135, 137	Variable third body reaction rates are used. The .chm file contains the average
	third body coefficients C_f and C_b , plus the names of the third body reactants and
	the corresponding third body efficiencies.

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Note: currently dissociation reactions with variable third body reaction rates (*ispec* values of 135 and 137) are only available for structured grids.

File Format. The information needed for each reaction is stored in the chemistry data (.chm) file, with the dissociation reactions listed first. I.e., reaction data is specified in the following order:

 $Information\ defining\ dissociation\ reaction\ 1$

. . .

Information defining dissociation reaction ndeq Information defining exchange reaction 1

. . .

Information defining exchange reaction nreg - ndeg

where nreq is the total number of reactions, and ndeq is the number of dissociation reactions.

First, for each of the dissociation reactions, the .chm file contains the following:

Contents of Record	Columns	Format
Record 1 (for all ispec values)		
• Name of first reactant	1-5	a 5
• Name of second reactant (blank for dissociation)	9-13	a5
• Name of first product	17 - 21	a5
• Name of second product	25 - 29	a5
• Temperature exponent S_f in equation for k_f	33–44	e12.4
• D_f/K_B in equation for k_f	45 - 56	e12.4
• Average third body coefficient C_f in equation for k_f	57 - 68	e12.4
Record 2 (for all ispec values)		
• Temperature exponent S_b in equation for k_b	33-44	e12.4
• D_b/K_B in equation for k_b	45 - 56	e12.4
• Average third body coefficient C_b in equation for k_b	57-68	e12.4
$P_{coords} = 1 \text{ (for isomes} = 195 \text{ 197)}$		

Records 3+ (for ispec = 135, 137)

 These records specify the third body reactants and the corresponding third body efficiencies. Only the non-unity efficiencies have to be specified. The first record in this group must be

```
THIRD [BODY] [EFFICIENCY] nnot1
```

where *nnot1* is the number of third body reactants with a non-unity efficiency. (The brackets are used to indicate that the words BODY and EFFICIENCY are optional.) The names of the third body reactants and the corresponding efficiencies are specified in the next *nnot1* records, in free format, as follows:

```
species(1) efficiency(1)
...
species(nnot1) efficiency(nnot1)
```

Following the data for the dissociation reactions, the .chm file contains the data for each of the exchange reactions. If ispec = 137, this section begins with an optional line containing either the word GENERAL or GLOBAL. I.e.,

[GENERAL | GLOBAL]

If this line is not present, or if *ispec* is not 137, only elementary exchange reactions with two reactants and two products may be used, and the data for each reaction is specified as follows:

Contents of Record	Columns	Format
Record 1		
• Name of first reactant	1-5	a5
• Name of second reactant	9-13	a 5
• Name of first product	17 - 21	a5
• Name of second product	25 – 29	a 5
• Temperature exponent S_f in equation for k_f	33 – 44	e12.4
• D_f/K_B in equation for k_f	45 - 56	e12.4
• Coefficient C_f in equation for k_f	57 - 68	e12.4
Record 2		
• Temperature exponent S_b in equation for k_b	33 – 44	e12.4
• D_b/K_B in equation for k_b	45 - 56	e12.4
• Coefficient C_b in equation for k_b	57–68	e12.4

If ispec = 137, and GENERAL or GLOBAL is specified, general or global exchange reactions with up to three reactants and three products are used, and the data for each reaction is specified as follows:

Contents of Record	Columns	Format
Record 1		
• Number of molecules and name of first reactant	1 - 15	f5.1,a10
• Number of molecules and name of second reactant	16 – 30	f5.1,a10
• Number of molecules and name of third reactant	31 – 45	f5.1,a10
• Number of molecules and name of first product	46 – 60	f5.1,a10
• Number of molecules and name of second product	61 - 75	f5.1,a10
• Number of molecules and name of third product	76 – 90	f5.1,a10
Record 2		
• Temperature exponent S_f in equation for k_f	3-14	g12.4
• D_f/K_B in equation for k_f	17-28	g12.4
• Coefficient C_f in equation for k_f	31 – 42	g12.4
Record 2+ (if GENERAL was specified)		
• Temperature exponent S_b in equation for k_b	3-14	g12.4
• D_b/K_B in equation for k_b	17 - 28	g12.4
• Coefficient C_b in equation for k_b	31–42	g12.4

 $Continued\ on\ next\ page$

(Continued from previous page)

Contents of Record

Columns Format

Record 3

• This section specifies the concentration exponents a-f in the forward and backward reaction rate equations. It starts with a line that must contain either the word STOICHIOMETRIC or EXPONENTS. I.e.,

{STOICHIOMETRIC | EXPONENTS}

If STOICHIOMETRIC is specified, the exponents a through f are set equal to the stoichiometric coefficients c_1 through c_6 , respectively.

If EXPONENTS is specified, the exponents a through f are specified by the following line, in free format, as follows:

a b c d e f

All six values must be specified. Note that since global exchange reactions are forward only, when <code>GLOBAL</code> is specified the exponents d through f aren't actually used, and may be specified as 0.0.

With Format 2, the data is read in subroutine frtin3, and the rates are computed in ratesbe.

Example

FINITE RATE COEFFICIENTS FROM EVAN & SCHEXNAYDER - CONVERTED TO EQUILIBRIUM CONSTANT FORM

	8,4	NREQ,NI TFRMIN	ΣEQ				
02	1.	0	1. 0	1.	-1.0	5.9340E+04	7.20E+18
					-1.0	0.0	4.00E+17
H2	1.	H	1. H	1.	-1.0	5.1987E+04	5.50E+18
					-1.0	0.0	1.80E+18
H20	1.	OH	1. H	1.	-1.5	5.9386E+04	5.20E+21
					-1.5	0.0	4.40E+20
OH	1.	0	1. H	1.	-1.0	5.0830E+04	8.50E+18
					-1.0	0.0	7.10E+18
02	1. H	1. OH	1. 0	1.	0.0	8.4550E+03	2.20E+14
					0.0	0.0	1.50E+13
H2	1. 0	1. OH	1. H	1.	0.0	5.5860E+03	7.50E+13
					0.0	4.4290E+03	3.00E+13
H20	1. 0	1. OH	1. OH	1.	0.0	9.0590E+03	5.80E+13
					0.0	5.0300E+02	5.30E+12
H20	1. H	1. OH	1. H2	1.	0.0	1.0116E+04	8.40E+13
					0.0	2.6000E+03	2.00E+13

Format 3; ispec = 120, 121

[Note - This format is considered obsolete, and may be removed in the future. The same capability is available using Format 2 with ispec = 137.]

This option is only for global 1- or 2-reaction chemistry, and only for forward reactions. It is a quick method for simulating detonation problems, for example, in which reactions proceed only forward to completion. It allows dissociation and exchange reactions of the form

$$c_1 S_1 \rightarrow c_3 S_3 + c_4 S_4$$

 $c_1 S_1 + c_2 S_2 \rightarrow c_3 S_3 + c_4 S_4$

Rate Equations. For dissociation reactions, the reaction rate equation is of the form

$$r = k[S_1]^a$$

For exchange reactions, the reaction rate equation is of the form

$$r = k[S_1]^a[S_2]^b$$

Rate Coefficient. The reaction rate coefficient k is computed using the equation

$$k = CT^S e^{-D/(K_B T)}$$

where T is the temperature in K, the ratio D/K_B is in K, and K_B is the Boltzmann constant. The reaction rate coefficient for the j'th reaction, C_j , has units $(\text{cm}^3/\text{g-mol})^{O_j-1}(\text{sec-K}^S)^{-1}$, where O_j is the order of the reaction (i.e., the total number of moles of reactants).

Eddy Dissipation Concept. For ispec = 121, the eddy dissipation concept may be used, as described above for Format 2 with ispec = 137.

File Format. The information needed for each reaction is stored in the chemistry data (.chm) file in the following order:

Information defining reaction 1

. . .

Information defining reaction nreq

where *nreq* is the total number of reactions.

The data to be specified in the .chm file is described in detail in the following table.

Contents of Record	Columns	Format
Record 1		
• Name and number of molecules of first reactant	1-8	a5,f3.1
• Name and number of molecules of second reactant (blank for dissociation)	9–16	a5,f3.1
• Name and number of molecules of first product	17 – 24	a5,f3.1

Continued on next page

(Continued from previous page)

Contents of Record	Columns	Format	
Name and number of molecules of second product	25-32	a5,f3.1	
ullet Temperature exponent S in equation for k	33 – 44	e12.4	
• D/K_B in equation for k	45 – 56	e12.4	
Record 2			
• Place holders for future use. Leave blank.			
Record 3			
ullet Concentration exponents a and b in the reaction rate equation	1-16	2f8.3	
ullet Coefficient C in equation for k	25 – 36	e12.4	

With Format 3, the data is read in subroutine frtin2, and the rates are computed in ratesf (for ispec = 120) or ratesg (ispec = 121).

Example

FINITE RATE COEFFICIENTS
WESTBROOK-DREYER GLOBAL REACTION MODEL

7.11.4 Transport Properties

This section of the chemistry data file contains the information used to compute the laminar viscosity and thermal conductivity. The formulation is based on Wilke's law, with either Sutherland's law or a NASA four-coefficient formula used individually for each species. Different constants or coefficients may be specified for different temperature ranges.

For Sutherland's law, the equations are of the following form:

$$\frac{\mu}{\mu_0} = \left(\frac{T}{T_{0\mu}}\right)^{\frac{3}{2}} \frac{T_{0\mu} + S_{\mu}}{T + S_{\mu}} \tag{1}$$

$$\frac{\kappa}{\kappa_0} = \left(\frac{T}{T_{0\kappa}}\right)^{\frac{3}{2}} \frac{T_{0\kappa} + S_{\kappa}}{T + S_{\kappa}} \tag{2}$$

For the NASA formula, the equations are (Svehla, 1995):

$$\ln \mu_n = A_\mu \ln T + \frac{B_\mu}{T} + \frac{C_\mu}{T^2} + D_\mu \tag{3}$$

$$\ln k_n = A_k \ln T + \frac{B_k}{T} + \frac{C_k}{T^2} + D_k \tag{4}$$

The data is read from the chemistry data file, in the following form:

TRANSPORT COEFFICIENTS [NASA]

Title, line 1
Title, line 2
Information for species 1
...

Information for species ns

The section title TRANSPORT COEFFICIENTS [NASA] must immediately follow the last data line in the "Reaction Rates" section, with no blank lines in between. If the optional word NASA is in the section title, the NASA formulas are used to compute the viscosity and thermal conductivity; otherwise Sutherland's law is used.

When Sutherland's law is used, the information for each species is stored in the .chm file as described in the following table. The reference viscosity μ_0 is in millipoise; the reference conductivity k_0 is in BTU/(hour-ft-°R); the reference temperatures and temperature offsets are in °R; and the beginning and ending temperatures for each curve are in K. Note that records 3 and 4 are repeated, in pairs, for each temperature range after the first. I.e., the Sutherland's law constants for both μ and k are listed for the first curve, followed by the constants for both μ and k for the second curve, etc.

Contents of Record	Columns	Format
Record 1		
• Name of species	1-8	a8
• Number of curves (i.e., temperature ranges) defining μ and k	11 - 15	i5
• Minimum and maximum temperature for first curve	16 - 35	2f10.3
• Reference viscosity μ_0 for first curve	37 - 48	e12.4
• Reference temperature $T_{0\mu}$ for first curve	49 – 60	e12.4
• Reference temperature offset S_{μ} for first curve	61 - 72	e12.4
Record 2		
• Reference conductivity k_0 for first curve	37 - 48	e12.4
• Reference temperature T_{0k} for first curve	49 – 60	e12.4
• Reference temperature offset S_k for first curve	61 - 72	e12.4
Record 3		
• Minimum and maximum temperature for curve	16 – 35	2f10.3
• Reference viscosity μ_0 for curve	37 - 48	e12.4
• Reference temperature $T_{0\mu}$ for curve	49 – 60	e12.4
• Reference temperature offset S_{μ} for curve	61 - 72	e12.4
Record 4		
• Reference conductivity k_0 for curve	37 - 48	e12.4
• Reference temperature T_{0k} for curve	49–60	e12.4
• Reference temperature offset S_k for curve	61–72	e12.4

When the NASA formula is used, the format for the information in the .chm file is as described in the following table. With the NASA formula, the viscosity μ is in micropoise, the thermal conductivity k is in microwatts/(cm-K), and the beginning and ending temperatures for each curve are in K. Note that record 2 is repeated for each viscosity temperature range, followed by record 3 repeated for each thermal conductivity temperature range. I.e., the coefficients for all the μ curves

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are listed first, followed by the coefficients for all the k curves.

Contents of Record	Columns	Format
Record 1		_
• Name of species	1 - 16	a16
• The letter "V"	35	a1
\bullet Number of curves (i.e., temperature ranges) defining μ	36	i1
• The letter "C"	37	a1
\bullet Number of curves (i.e., temperature ranges) defining k	38	i1
Records 2+		
• The letter "V"	2	a1
• Minimum and maximum temperature for curve	3-20	2e9.2
• Coefficients A_{μ} , B_{μ} , C_{μ} , and D_{μ} for curve	21 – 80	4e15.8
Records 3+		
• The letter "C"	2	a1
• Minimum and maximum temperature for curve	3-20	2e9.2
• Coefficients A_k , B_k , C_k , and D_k for curve	21-80	4e15.8

If the effective binary diffusion model is being used (specified by the the DIFFUSION EFFECTIVE-BINARY keyword in the CHEMISTRY keyword block), the Lennard-Jones parameters used to compute the effective binary diffusivity for each species must also be included in the .chm file, following the TRANSPORT COEFFICIENTS data described above. Values for these parameters for a variety of species may be found in the literature (e.g., Bird, Stewart, and Lightfoot (1960).)

The data is read in the following form:

LENNARD-JONES PARAMETERS

Title, line 1

Title, line 2

Parameters for species 1

. . .

Parameters for species ns

The title LENNARD-JONES PARAMETERS must immediately follow the preceding data line, with no blank lines in between.

For each species, the data is read using the format described below. The parameter σ is in angstroms, and ϵ/K_B is in degrees Kelvin.

Contents of Record	Columns	Format
• Name of species	1-8	a8
• Lennard-Jones parameter σ	37 - 48	f12.4
• Lennard-Jones parameter ϵ/K_B	49 – 60	f12.4

Example

TRANSPORT COEFFICIENTS

02	1	300.000	15000.000	1.9190E-01	4.9160E+02	2.5000E+02
				1.4190E-02	4.9160E+02	4.0000E+02
NO	1	300.000	15000.000	1.3700E-01	4.9160E+02	4.0000E+02
				8.4070E-03	4.9160E+02	4.0000E+03
0	1	300.000	15000.000	1.7030E-01	7.5000E+02	1.5500E+03
				1.0360E-02	4.9160E+02	2.3000E+03
N	1	300.000	15000.000	1.7890E-01	4.9160E+02	2.3000E+02
				6.8900E-02	4.9160E+02	2.3030E+02
N2	1	300.000	15000.000	1.6630E-01	4.9160E+02	1.9200E+02
				1.4000E-02	4.9160E+02	3.0000E+02

Keywords: CHEMISTRY

7.12 Memory Log File (memdebug.lis)

Fortran unit number 97

A $memory\ log\ file$, always named memdebug.lis, may be created using <code>DEBUG</code> option 65 to track memory allocation/deallocation requests.

Keywords: DEBUG 65

7.13 Reserved Files

Unit numbers 15, 32, and 55 are reserved for proprietary features, and should not be used by Wind-US developers.

8 Scripts

8.1 wind — Run Wind-US

The usual procedure for running Wind-US is via a command-line script called *wind*. With this approach, the specific executable to be run (e.g., production version, alpha version, etc.), the names of the required input and output files, etc., may be defaulted, specified on the command line, or entered as responses to prompts.¹⁶

The syntax used when invoking wind is:

The various command-line options are described below. Note that all are optional. Several use the syntax -(no)option, indicating that they may be specified as either -option or -nooption. In addition to specifying values on the command line, several of the options may also be specified using environment variables. Values specified on the command line will override values set by environment variables.

	`	•	
- (no l	bat	:ch

If -batch is specified, run in "batch" mode, with all necessary input specified via command-line options. The user must specify the executable to be run using the -program option; all the input/output file names using the -dat, -list, -grid, and -flow options; the system run queue using the -runque option. The user must also specify either the -runinplace option, or the directory to run in using the -runroot option. In addition, if a multi-processing control (.mpc) file exists, either the -parallel or -noparallel option must be specified. And, if an .mpc file exists and MPI message passing is being used, the number of zones must be specified using the -nzones option. If any of these items are missing, an error message will be displayed and execution will stop.

The default is -nobatch, allowing the necessary input to be specified in response to prompts.

-begtime time

The beginning time for a standard Unix *at/batch* job (i.e., when using -runque AT_QUE).

-(no)bg

Jobs in the interactive queue normally run in the foreground if the output is being displayed on the screen, and in the background if the output is being sent to the *.lis* file. If -nobg is specified, interactive

 $^{^{16}}$ The wind script is actually a simple wrapper, that passes its arguments through to wind.pl, which in turn is a symbolic link to a Perl script named solver.pl. A job file (with the suffix .job.pl) is created on the fly, which is the actual script submitted to the queue specified by the user. This is all normally transparent to users, however.

the .lis file.

Root location of the Wind-US executable on the system on which it -binroot directory will be running. Environment variables: WIND_BINROOT, CFDROOT_REM. In order of preference, the default root location is set to the value of the environment variable WIND_BINROOT (if defined), then CFDROOT_REM (if defined), then CFDROOT. -cfdrootrem directory The CFDROOT directory on the system on which the Wind-US executable will be running. The default is the same as the CFDROOT directory on the local machine. Environment variable: CFDROOT_REM. -(no)cgesave By default, when Wind-US is being run in a remote directory (i.e., in master mode, or without -runinplace), the common edge (.cge) file used for unstructured grids is saved in the job submittal directory at the end of the run. Specifying -nocgesave indicates that the .cge file is not to be saved. -(no)clSpecifying -cl indicates that a cluster machine PVM environment (e.g., a Linux cluster) is to be assumed. The default is -nocl. Environment variable: WIND_CL. Set to YES for a cluster, NO otherwise. -cleanup Perform clean-up functions only, copying files to the run directory if necessary, and removing temporary files, then exit. These clean-up functions are normally done only during normal job termination, or in the event of an error during the run. -dat dat fileThe input data (.dat) file, entered without the extension (e.g., wing, not wing.dat). If -debug is specified, Wind-US will run in debug mode, using the de--(no)debug bugger specified by the -debugger option. The job will automatically be run interactively, and the -runinplace option will be set. The default is -nodebug. Environment variable: WIND_TEST_MODE. Set to DEBUG for debug mode, NODEBUG otherwise. -debugger debugger Define the debug package to be used when running in debug mode. The default depends on the computer system being used. Environment variable: WIND_DEBUGGER. Create a file named *done* at the end of the run containing a brief job -done status message. The status message will be "Done" for normal termination, "Execution Interrupted" for an interrupt during job execution, or "Error On Exit" for an error during job termination. -endtime timeEnding time for a Unix at/batch job, or time to run in seconds for a job running under the basic QSUB, PBS, or LSF queueing system. For the PBS and LSF queueing systems, some additional time will

jobs will be run in the foreground, even if the output is being sent to

be added for termination processing (i.e., updating the flow (.cfl) file, terminating worker processes, etc.) Also see the -walltime option.

-flow cflfile

The flow (.cfl) output file, entered without the extension (e.g., wing, not wing.cfl). If the file exists, the solution will restart using the conditions in the file as initial conditions.

-genpost time

Create/update GENPLOT files containing convergence information every time seconds. If <code>-genpost</code> is specified without a time, the value specified with <code>-waittime</code> (or its default value) is used. Note that this option requires use of the <code>POSTPROC</code> keyword block in the input data (.dat) file. The <code>-genpost</code> option may not be used for jobs in the interactive queue with output sent to the screen.

Also see the -loads option for a quick, but less general, way to create a GENPLOT file containing convergence information.

-grid cgdfile

The grid (.cgd) input file, entered without the extension (e.g., wing, not wing.cgd).

-grpcharge number

Group charge number for jobs run under the PBS queueing system on Cray machines.

-help

Display the list of command-line options, and quit, ignoring any other options.

-list *lisfile*

The list output (.lis) file, entered without the extension (e.g., wing, not wing.lis). For interactive jobs, set lisfile to "screen" to display the .lis file at the terminal.

-loads time

Create/update a GENPLOT file containing total forces and moments, and the L2 norm of the Navier-Stokes residual, every time seconds. The file name will be lisbase.loads.gen, where "lisbase" is the base name of the .lis file. If -loads is specified without a time, the value specified with -waittime (or its default value) is used. The LOADS keyword block must be used to write the desired total loads data (i.e., using TOTALS with the PRINT keyword) into the .lis file. The -loads option may not be used for jobs in the interactive queue with output sent to the screen.

Also see the **-genpost** option for a more general way to create GEN-PLOT files containing convergence information.

Due to a script bug, the -loads option will not work with versions of the Wind Scripts older than 2.0.82. This corresponds to Wind-US versions obtained before the afternoon of 14 Mar 2008.

-memory number

Amount of run-time memory (in megabytes) to request for a job running under the PBS queueing system.

-(no)mp

Specifying -mp indicates that a multi-processor machine (i.e., a single machine with multiple processors, like the CRAY or SGI Origin-2000) is being used, with either PVM or MPI message passing. The default is -nomp.

Environment variable: $WIND_MP$. Set to YES for a multi-processor machine, NO otherwise.

-mpiver version

MPI distribution being used when running in parallel with MPI message passing. *version* must be one of OPENMPI, MPICH1, MPICH2, INTEL, SGI. The default is MPICH2.

Environment variable: WIND_MP_VER.

-mpmode mode

Message passing mode to be used when running in parallel: either PVM or MPI. Note that the pre-compiled executables available through IVMS were built without MPI message passing. To use MPI message passing, MPI must be pre-installed on your system(s) (unlike PVM, MPI is not distributed with Wind-US), and you'll need executables that includes links to the MPI library. See the section "Installing the Build Distribution" in the Wind-US Installation Guide for instructions on creating the executable. The default message passing mode is PVM.

Environment variable: WIND_MP_MODE.

-ncpu number

Number of CPUs to request for a job running under the PBS or LSF queueing system.

-nice_val number

The "nice" value (a number from 1 to 20) to use with the Unix *nice* command. Larger values cause the job to run at a lower CPU scheduling priority to lessen the impact on other jobs on the system. For interactive jobs (-runque REAL), the value is automatically set to 2.

Environment variable: WIND_NICE_VAL.

-norsh

No remote shell (i.e., rsh or ssh) capability is available. This option only applies when MPI message passing is being used on distributed systems; all systems must be of the same type, and the Wind-US executable must be in the same location on each system. Parallel runs with PVM message passing on distributed systems and clusters require a remote shell capability.

Environment variable: WIND_USE_SSH. Set to NONE for no remote shell capability.

-not_nice

Run without using the Unix *nice* command. This option only applies to serial jobs, and to worker tasks in parallel jobs that are being run on a different system than the master. For parallel jobs, the master task, and worker tasks being run on the same system as the master, automatically run at top priority (i.e., without using *nice*).

Environment variable: WIND_NOT_NICE. Set to YES to run without the *nice* command, NO otherwise.

-nzones number

Number of zones, used in MPI message passing mode. If not specified, the utility mpigetnzone will be used to get the number of zones from the common grid (.cgd) file. If mpigetnzone is not installed, the user will be prompted for the number of zones.

-(no)parallel

Specifying -parallel indicates that the job is to be run in parallel mode, while -noparallel indicates serial mode. Parallel mode requires a multi-processing control (.mpc) file. If -parallel is specified, but an .mpc file doesn't exist, the user will be asked if serial mode should be used. If neither -parallel nor -noparallel is specified,

and an .mpc file exists, the user will be asked if parallel mode should be used.

-program windname

The name of the executable to be used, without the .exe extension.

-queue_name name

In the queueing system being used, as specified using the **-runque** option, the name of the specific queue in which the job is to run. The form of the *name* parameter depends on the queueing system.

-remoterun

Bypass some of the built-in error checking for valid path names preventing error messages when running on a remote system. The CFD-ROOT directory on the remote machine should be specified using the -cfdrootrem option, and the root directory under which the Wind-US executable is to be run on the remote system should be specified using the -runroot option.

-runinplace

Run in place, i.e., in the directory in which the *wind* script was invoked. This option is automatically selected if master mode is being used, or if running in debug mode. Otherwise, the default is to run in the directory specified using **-runroot**.

Environment variable: WIND_RUNINPLACE. Set to YES to run in place, NO otherwise.

-runque queue

Queueing system in which the Wind-US executable is to run. The parameter *queue* may be REAL for a real-time interactive job; AT_QUE to run in a Unix *at* or *batch* queue; QSUB_QUE to run in a basic QSUB queue; QSUB_PBS_QUE to run in a QSUB queue using the PBS batch system; or BSUB_QUE to run in a BSUB queue using the LSF batch system. If the specified queueing system doesn't exist on the system being used, the job will stop.

Environment variable: WIND_RUN_QUE.

-runroot directory

Root directory under which the Wind-US executable is to be run when -runinplace is not specified. The full name will be *directory/userid/basename.scr*, where *userid* is your userid and *basename* is the base name of your .dat file. The default directory, if -runroot is not specified, is /tmp.

Environment variable: WIND_RUNROOT.

-(no)save_core

Specifying -save_core sets the allowed core file size equal to the maximum possible value. The default is -nosave_core, which makes no change to the allowed size. This option only applies to Linux systems.

-solver solver

The solver to be run. The default, and only meaningful value for running Wind-US, is the base name of the top level Perl script (i.e., wind, from wind.pl).

-test

Sets the root location of the Wind-US executable to the value of the environment variable WIND_DEV, allowing a "test" version to be used without having to change your CFDROOT environment variable. (More specifically, it's set to SOLVER_DEV, where SOLVER is the solver being run, specified using -solver, converted to upper case.) This option overrides a value specified using -binroot.

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-tmpdir directory The directory for storing temporary files during a run. The default

is defined by the script <code>pvmgettmp.pl</code>, which sets it to the value of the environment variable <code>PVM_TEMP</code> (if defined); otherwise it's set to the first of the following directories that exists: <code>/lscratch, /scratch*</code>,

/data/local, /tmp.

Environment variable: TEMPDIR.

-user_name name The username to be used for Windows Scheduler. The default is the

current user. (Note, though, that Wind-US is not officially supported

under Windows.)

-user_passwd passwd The user password to be used for Windows Scheduler. There is no

default value. (Note, though, that Wind-US is not officially supported

under Windows.)

-(no)usessh When -usessh is specified, ssh/scp remote shell/copy commands will

be used when copying files between the master and workers when running in parallel mode on distributed systems or clusters. The default is to use rsh/rcp. For more details see the discussion of remote shell

commands for parallel processing in Section 9.3.

Environment variable: WIND_USE_SSH. Set to YES to use ssh/scp, NO

to use rsh/rcp.

-version Display the specific version numbers for the Wind-US executable, and

the CFD, ADF, and PVM libraries it's linked with, and immediately

exit.

-waittime time Wait time seconds between run status checks. The default value, if

-waittime is not specified, is 10 seconds. If -waittime is specified,

but with a value less than 1, the value is reset to 30.

-walltime time Total wall clock time in seconds for a job running under the PBS or

LSF queueing system. This includes the length of time for termination processing (i.e., updating the flow (.cfl) file, terminating worker

processes, etc.) Also see the -endtime option.

8.2 wind_post — Perform Post-Processing

At the end of a non-debug job that completes successfully, if a script named wind_post exists in the current directory, and if the job is not being run interactively, the wind_post script is invoked, as follows.

wind_post datname lisname cgdname cflname iter run_command

The six arguments are defined as:

datname The base name (i.e., without the three-letter extension) of the input data (.dat)

and job files

lisname The base name of the list output (.lis) file

cgdname The base name of the grid (.cqd) file. (If CGNS files are being used, note that

this is the base name of the file itself, not the name of the CGNSBase_t node in

the file.)

cflname The base name of the flow (.cfl) file

iter The "job iteration" number. This is a count of the number of jobs, not time

step iterations. It is initialized to zero, and incremented by one before each invocation of *wind_post*. Note that the value of *iter* is not saved between runs. The next invocation of the top-level *wind* script will re-initialize the value to

zero.

run_command The command used to submit the job

The wind_post script itself must be supplied by the user. Users with some experience in writing shell scripts may use this feature to automatically post-process results at the end of a job, or to save and/or process interim results from a lengthy calculation and submit a new job.¹⁷

An example wind_post Bourne shell script called wind_post.sh is supplied with the Wind-US application distribution in the directory wind/bin. (There's also an example C shell version called wind_post.csh.) The example script resubmits the job with a different input file for each job iteration. It looks for an input data file named datname.dat.nextiter, where datname is the base name of the .dat file and nextiter is the next job iteration number (i.e., one more that the value of the input argument iter). The new input data file, if it exists, is copied over the original input data file and the job file is resubmitted.

A typical method for using this capability would be to create files named, for example, test.dat.1, test.dat.2, test.dat.3, etc. Then, copy test.dat.1 to test.dat and submit the job as usual. By keeping the initial input file in test.dat.1 and copying it to test.dat, the original initial input file is preserved.

After the initial job completes, $wind_post$ will be called with iter = 1. The next job iteration number is thus 2. The file test.dat.2 will be copied into test.dat, and the job will be resubmitted, restarting Wind-US where it left off. If the file test.dat.nextiter doesn't exist, all the input data files have been used. The job file is then removed, and the run ends.

Note the distinction between a run and a job, and the value of the job iteration number. By run, we mean here a single invocation by the user of the wind script. Within that run, multiple jobs may be submitted using the wind_post feature, each with a different, automatically-incremented job iteration number. As noted above, the job iteration number is not saved between runs. The next time the wind script is invoked, starting a new run, the job iteration number will be re-initialized to zero. If it's necessary to save the value between runs, the user-supplied wind_post script should save it to a file, to be read during the next run.

The example wind_post script does not do any post-processing, but that can easily be added. For example, CFPOST could be run, with input redirected from a command file, to create plot files or reports. Or, an interim flow (.cfl) file could be saved by copying it into a file with the iter variable as part of the file name.

8.3 windver — Get Wind-US Version Number

The alpha version of Wind-US, and to a lesser extent the beta version, change quite frequently. Even the released production version is sometimes modified, as a result of bug fixes. Changes are summarized in the "Source History" file, accessed through the Internet Version Management System (IVMS).

The *windver* script may be used to get the complete version number of a Wind-US executable, and its supporting libraries. For example,

 $^{^{17}}$ Note that a similar capability may be invoked using the SPAWN keyword in Wind-US's input data file.

```
% windver

***** Print Version numbers for Wind *****

--Solver bin directory set to /usr/local/wind

Select the desired version

0: Exit wind
1: Wind 5.0
2: Wind-US 1.0
3: Wind-US 2.0

Enter number or name of executable......[3]: 3

Version numbers for /usr/local/wind/LINUX32-GLIBC2.3/XEON/bin/Wind-US2.exe

WINDUS - Version 2.215 (Last changed 2007/08/28 14:00:42)
LIBCFD - Version 2.167 (Last changed 2007/08/22 23:44:18)
LIBADF - Version 2.0.15.1 (Last changed 2007/05/31 21:58:08)
PVM - Version 3.4.5
```

windver lists the various executables available, and prompts the user to select one of them. It then prints the complete version number, plus the date it was created, for Wind-US, the CFD library (common file processing routines, system-dependent routines, and utility routines), the ADF library (ADF routines from the CGNS project), and the PVM library (PVM routines used for parallel processing). In the above example, Wind-US 2.0 is actually Version 2.215, created on Aug 28, 2007.

By default, the windver script looks for executables in the directory specified by the environment variable WIND_BINROOT. If there is no environment variable of that name, windver will look in CFDROOT. To look in some other location, instead of using windver, use the regular wind script with the -binroot and -version options. E.g., if the executables are located in the appropriate directories (i.e., in subdirectories corresponding to the SYSTEM and SYSTEM_CPU environment variables) below /home/wind_test/, you would do:

wind -binroot /home/wind_test -version

8.4 windrun — Quick Wind-US run

The windrun script may be used to quickly set up and start a Wind-US run. The run will be in interactive mode, with the output sent to a .lis file. Internally windrun invokes the wind script with the options -nobg and -runque REAL, and with other wind script options determined by the windrun options.

The syntax used when invoking windrun is:

The command-line arguments and options are described below. Note that except for *casename*, all are optional.

casename

The prefix for the Wind-US input/output files. All files must have the same prefix (e.g., case1.cgd, case1.cfl, etc.)

-dev | -cfd

Specifies where to look for the Wind-US executable. If <code>-dev</code> is specified, look under the directory defined by the <code>WIND_DEV</code> environment variable. If <code>-cfd</code> is specified, look under the directory defined by the <code>CFDROOT</code> environment variable. The default is to look under the directory defined by the <code>WIND_DEV</code> environment variable if the <code>WIND_DEV</code> environment variable exists, and under the directory defined by the <code>CFDROOT</code> environment variable if the <code>WIND_DEV</code> environment variable does not exist.

windname

The name (without the .exe extension) of the Wind-US executable to be used (e.g., windalpha, not windalpha.exe).

-serial | -parallel

If -serial is specified, run in serial mode. If -parallel is specified, run in parallel mode on a multiprocessor system (i.e., using the -mp option to the wind script), if an .mpc file is present. The default is to run in serial mode.

-new | -restart

If -new is specified, run a new case from scratch. Any existing .cfl, .tda, .cth, .lis, .job, .cge, and .cft files with the prefix casename, and any memdebug.lis file, in the current directory will be erased before starting the Wind-US run. If -restart is specified, start from the the existing solution, if one exists. The default is to run a new case from scratch.

-runlocal | -runtmp

If -runlocal is specified, use the -runinplace option to the wind script to run in the current directory. If -runtmp is specified, use the options -remoterun -runroot \$TMPDIR to run in a temporary directory below the location specified by the TMPDIR environment variable. The default is to run in the current directory.

-help

Display a summary on using windrun.

8.5 windmp — Run on Multi-Processor

The windmp script may be used as a shortcut when running Wind-US on a multi-processor machine (i.e., a single machine with multiple processors, like the CRAY or SGI Origin-2000). It simply invokes the wind script with the options -mp, -runinplace, and -not_nice.

8.6 runtest, runtestsuite — Run Wind-US Test Case(s)

runtest — Run Wind-US test cases

runtest is a utility for running Wind-US test cases. The syntax used when invoking runtest is:

The base name of the case to be run is given by the specified *casename*. runtest will look for *casename.dat* and *casename.cgd* input files in the current directory. In addition, runtest will

recursively descend into any sub-directories and look for input files named *subdir.dat* and *subdir.cgd*, where *subdir* is the sub-directory name.

For example, you could set up a test case named *case1*, with input files *case1.dat* and *case1.cgd*. In the directory containing these files, you could have sub-directories named (say) *run1*, *run2*, etc., containing variations (e.g., different input options) on the original case. *runtest* will *cd* into the sub-directory *run1*, and look for input files named *run1.dat* and *run1.cgd*. Similarly for the sub-directory *run2*, etc. All such cases that are found will be run, in the directory containing the input files.

If a file named *casename.ic.cfl* (or *subdir.ic.cfl* if in a sub-directory, as described above) exists in the run directory, it will be copied as the file *casename.cfl* (or *subdir.cfl*) and used as initial conditions. Otherwise, a new case will be run from scratch. In either situation, any existing output files will be overwritten.

If multiple .dat files are found named casename.1.dat, casename.2.dat, etc. (or subdir.1.dat, etc.), multiple runs will be made using the successive .dat files to restart the solution.

If the specified case has been run previously, any convergence measures stored in the .lis file (i.e., residuals, integrated flow properties, etc.) will be compared with the values from the existing baseline results using diff, and the output will be written to the file basedir/test.log. In addition, if a file named delta.com exists in the directory with the Wind-US input files, it's assumed to contain CFPOST commands. CFPOST is run, with its standard input redirected from delta.com, and its standard output appended to the file basedir/test.log.

The -baseline option may be used to replace an existing baseline solution with the current one, bypassing the comparison of results.

The command-line arguments and options are described below. Note that except for basedir and casename, all are optional.

basedir The directory where the log file of the tests is to reside.

The prefix for the Wind-US input/output files. All files must have the

same prefix (e.g., case1.cgd, case1.cfl, etc.)

-dev | -cfd Specifies where to look for the Wind-US executable. If -dev is specified, look under the directory defined by the WIND_DEV environment

variable. If -cfd is specified, look under the directory defined by the CFDROOT environment variable. The default is to look under the directory defined by the WIND_DEV environment variable if the WIND_DEV environment variable exists, and under the directory defined by the CFDROOT environment variable if the WIND_DEV environment variable

does not exist.

windname The name (without the .exe extension) of the Wind-US executable to

be used (e.g., windalpha, not windalpha.exe). If no name is specified,

the default Wind-US version will be used.

-serial | -parallel | If -serial is specified, run in serial mode. If -parallel is specified, run in parallel mode on a multiprocessor system (i.e., using the -mp

option to the *wind* script), if an .mpc file is present. The default is to

run in serial mode.

-runlocal | -runtmp If -runlocal is specified, use the -runinplace option to the *wind* script to run in the directory where the input files are found. If -

runtmp is specified, use the options -remoterun -runroot \$TMPDIR to run in a temporary directory below the location specified by the

 ${\tt TMPDIR}$ environment variable. The default is to run in the current

directory.

-baseline Replace the saved baseline solution, used for comparison with results

from future runs, with the current solution.

-help Display a summary on using runtest.

runtestsuite — Run a series of Wind-US test cases

runtestsuite is a wrapper for runtest that may be used to automate the running of a series of test cases. It assumes that the cases to be run are stored in subdirectories of the current directory, with the subdirectory names the same as the case names (i.e., the subdirectory names are used for the casename argument when invoking the runtest script).

The syntax used when invoking *runtestsuite* is:

runtestsuite [-dev | -cfd] [windname] [-serial | -parallel] [-baseline] [-help]

The command-line arguments and options are described below. Note that all are optional.

-dev | -cfd Specifies where to look for the Wind-US executable. If -dev is spec-

ified, look under the directory defined by the WIND_DEV environment variable. If -cfd is specified, look under the directory defined by the CFDROOT environment variable. The default is to look under the directory defined by the WIND_DEV environment variable if the WIND_DEV environment variable exists, and under the directory defined by the CFDROOT environment variable if the WIND_DEV environment variable

does not exist.

windname The name (without the .exe extension) of the Wind-US executable to

be used (e.g., windalpha, not windalpha.exe). If no name is specified,

the default Wind-US version will be used.

-serial | -parallel | If -serial is specified, run in serial mode. If -parallel is specified,

run in parallel mode on a multiprocessor system (i.e., using the -mp option to the wind script), if an .mpc file is present. The default is to

run in serial mode.

-baseline Replace the saved baseline solutions for each case, used for comparison

with results from future runs, with the current solutions.

-help Display a summary on using runtestsuite.

9 Parallel Processing

The time required to compute a solution can be reduced dramatically by using the parallel processing capability of Wind-US, with multiple zones being solved simultaneously. Jobs may be run in parallel mode on a multi-processor system (i.e., a single system with multiple CPUs), on a workstation cluster (i.e., a collection of networked nodes designed for parallel computation, with NFS-mounted home directories), or on a collection of separate, possibly heterogeneous, distributed networked systems, with or without NFS-mounted home directories. Either PVM or MPI message passing may be used on multi-processor systems; only PVM is supported when running on a cluster or on distributed networked systems.

Running parallel jobs with Wind-US is remarkably simple. The only user requirement, beyond some initial system and account setup needed for communication, is the creation of a file listing the systems and/or number of processes to be used. The Wind-US scripts automatically take care of copying the necessary files and executables between the systems being used, starting and stopping the message passing software, and cleaning up at the end of the run.

9.1 Terminology

When operating in parallel processing mode, the system or node on which the job originates is called the master.¹⁸ The nodes used to perform the actual solution are called workers. The master distributes grid zones to participating workers. Each zone is solved in parallel with other zones on other workers. Boundary information is exchanged at the end of each cycle to propagate information throughout the computational domain. There may be fewer workers than zones to be computed, in which case, a worker will be assigned another zone when it finishes its current assignment. The user specifies the names of the participating workers via the multi-processing control (.mpc) file, described in Section 9.6.

9.2 Parallel-Capable Executables

At the time a Wind-US executable is built, it's linked with the appropriate message passing libraries (i.e., PVM, MPI, or both) that are required for parallel operation. The Wind-US software distributed by the NPARC Alliance includes PVM, but does not include MPI. If MPI is to be used, the MPI libraries and executable must be pre-installed on the multi-processor system being used.

The makefiles used to build the Wind-US executable include switches that specify the message passing software to be used. The default is PVM, and the executables available on IVMS are built for PVM only. If MPI is to be used, an executable must be built from the build distribution, with the appropriate makefile switches set. See the Wind-US Installation Guide for details.

If Wind-US will be run in parallel mode on a collection of heterogeneous networked systems, executables must be available for each of the system and CPU types that will be used, and stored on the originating system. The appropriate Wind-US and PVM executables will automatically be copied to the workers at the start of the run (and to the master when master mode is used), and removed when the run finishes. Details on the directory structure required for the executables on the originating system are presented in Section 9.4.

¹⁸Strictly speaking, the master is the system controlling the job, not necessarily the system used to submit the job. When "master mode" is used, a system different from the originating system may be specified as the master. See the description of the .mpc file #master directive in Section 9.6 for details.

The run scripts used with Wind-US are designed to use the PVM executables that are part of the Wind-US distribution. If PVM happens to be pre-installed on the system(s) being used, some PVM-related environment variables may already be defined, such as PVM_ROOT or PVM_HOME. (The environment variables may be listed by issuing the command setent, assuming the C shell or one of its variants is being used as the interactive shell.) In some cases, these have been found to conflict with the execution of Wind-US with its own version of PVM. In this case, the PVM-related environment variables should be unset before starting Wind-US. If the pre-installed version of PVM isn't needed for running other applications, a simple way to do this is to add a line like

unsetenv PVM_ROOT

in your shell startup file (i.e., .chsrc for the C shell).

9.3 Remote Shell Commands

To run in parallel mode on a cluster or on distributed networked systems, the user must of course have a valid account on each system. The user name on all the systems must be the same.

In addition, the master must be allowed to communicate with each worker, and vice versa, using remote shell commands, and without entering a password. By "remote shell commands" we mean either rsh (remsh on HP and Cray systems) and rcp, or ssh and scp. ¹⁹ If TCPD access control is installed, which remote shell commands are allowed is normally controlled at the system level by information in the files /etc/hosts.allow and /etc/hosts.deny.

The following two sections describe how to set up password-less communication between the master and workers for rsh/rcp and for ssh/scp. If master mode is used, the same procedure must be followed to set up password-less communication between the originating system and the master, as well as with each worker. Note that this only has to be done once for a given cluster or collection of distributed systems.

Important: Wind-US uses the Unix hostname command to determine a system's name. Thus, in the procedures described below, whenever a system name is specified the name to be used must be the same as the name returned by the hostname command on that system. For example, for a system with the fully-qualified name "workerbee.bigcompany.com", if hostname returns just the machine name "workerbee", then workerbee should be used for the system name. If hostname returns the fully-qualified name "workerbee.bigcompany.com", then workerbee.bigcompany.com should be used for the system name.

$9.3.1 \quad rsh/rcp$

To allow rsh and rcp to be used without entering a password, the host name of the master system must be in the file .rhosts in the user's home directory on the worker system, or in the system file /etc/hosts.equiv on the worker system, and vice versa. Note that this is required even if the master and worker are the same system. I.e., if the master is also being used as a worker, that system's name must be listed in the .rhosts or /etc/hosts.equiv file.

The *.rhosts* file is a text file containing a list of system names, and the userids on each of those systems, that are allowed to access the current host via *rsh* and *rcp*. The file should have its permissions set to rw----, so issue the following command after creating the file.

 $^{^{19}}$ Note that rsh and rcp are considered insecure, and many organizations, if not most, now require use of ssh and scp.

```
chmod 600 .rhosts
```

Once the *.rhosts* file has been created, it may be tested by issuing the following command from the system where the job will be submitted.

```
rsh worker-name ls -1
```

Things are functioning properly if the directory listing appears.

When rsh/rcp remote shell commands are used, the maximum number of hosts that may be specified in the .mpc file is limited to 25, due to timeout issues that can occur with rsh/rcp.

More information about .rhosts files may be found by entering man rhosts on most Unix systems.

9.3.2 ssh/scp

Setting up password-less communication using ssh and scp is a bit more complicated, but as noted earlier it only needs to be done once for a given cluster or collection of distributed systems.

When performing the following steps, when you use ssh to connect to a new system (i.e., one that you've never connected to using ssh, or one whose host key has been changed), you may get a message like

```
The authenticity of host 'system_name (133.11.217.42)' can't be established. RSA key fingerprint is ec:73:17:40:8d:c0:b5:96:76:27:6b:ce:f4:f9:96:73. Are you sure you want to continue connecting (yes/no)?
```

If so, respond with "yes" (without the quotes). This will add the host key for $system_name$ in your $.ssh/known_hosts$ file.

1. First, on the master, create private and public authentication keys by doing

```
ssh-keygen -t rsa
```

The option -t rsa means use RSA authentication. If your site uses DSA authentication, you should use the option -t dsa. Use the defaults (i.e., just hit Enter) when prompted for a file name and passphrase (i.e., use no passphrase, in order to allow password-less ssh connections).

This creates, in your /.ssh directory, the files id_rsa , containing your private authentication key, and $id_rsa.pub$, containing your public authentication key. Make sure the id_rsa file is only readable by you. I.e., in the .ssh directory, doing "ls -l id_rsa" should give something like:

```
-rw----- 1 userid userid 883 Jun 23 09:12 id_rsa
```

If it doesn't, do "chmod 600 id_rsa".

2. Still on the master, add the public authentication key to the file .ssh/authorized_keys in your /.ssh directory by doing:

```
cd .ssh
cat id_rsa.pub >> authorized_keys
```

- 3. If you're using a cluster or distributed systems with NFS-mounted home directories (i.e., your home directory physically resides on one node, and is NFS-mounted, or "shared", with the other nodes), do the following.
 - For each node, including the master, do

```
ssh node\_name ls
```

where node_name is the node name. If you get the "authenticity of host ... can't be established." message described earlier, respond with "yes". This will ensure that each node is listed in your .ssh/known_hosts file (or is already in the system-wide ssh_known_hosts file).

• In your .ssh directory, edit the authorized_keys file. There should already be a long line there for the master that was created in step 2, that looks something like

```
ssh-rsa public_key= userid@master
```

where *public_key* is a long string of characters containing your public authentication key, *userid* is your user ID, and *master* is the name of the master node. For each additional node, copy this line, and change the name *master* at the end of the line to the name of the node.

- 4. If you're using distributed systems with separate home directories on each system, do the following.
 - From the master, add the master's public authentication key to the file .ssh/authorized_keys on each worker. I.e, from the .ssh directory on the master, for each worker system do the following. (Here, and in the following instructions, replace "worker" with the name of the worker system.)

```
cat id_rsa.pub | ssh worker 'cat >> .ssh/authorized_keys'
```

• Log in to each worker system, and create private and public authentication keys on that system. I.e., from the master you could do the following.

```
ssh ital
ssh-keygen -t rsa
cd .ssh
cat id_rsa.pub >> authorized_keys
```

• On each worker, add that worker's public authentication key to the file .ssh/authorized_keys on the master. I.e., from the .ssh directory on each worker, do the following (where "master" is replaced with the name of the master system).

```
cat id_rsa.pub | ssh master 'cat >> .ssh/authorized_keys'
```

You should now be able to use ssh (and scp) from the master to a worker, and vice versa, without entering a password. To test this, on the master do

```
ssh worker ls -1
```

The contents of your home directory on the worker should be displayed. Similarly, on the worker do

```
ssh master ls -1
```

The contents of your home directory on the master should be displayed.

Note that if the master is also being used as a worker, you must also be able to use *ssh* locally. To test this, on the master do

```
ssh master ls -1
```

When running Wind-US, use the **-usessh** option to the *wind* script to specify that the executable and I/O files should be copied between the master and workers using *ssh*. I.e.,

```
wind -usessh
```

9.4 Directory Structure for Executables

The run scripts expect to find the Wind-US executable(s) (Wind-US2.exe, for Wind-US 2.0) and the PVM executables (pvm, pvmd3, and pvmgs) for particular systems and CPUs in specific locations below the CFDROOT directory on the originating system, corresponding to the values of the SYSTEM and SYSTEM_CPU environment variables for those systems.²⁰

As an example, suppose the systems being used are a mix of 32-bit Linux systems with XEON processors and *glibc* version 2.3 (i.e., SYSTEM = LINUX32-GLIBC2.3 and SYSTEM_CPU = XEON), 64-bit Linux systems with OPTERON processors and *glibc* version 2.3, and 64-bit SGI R14000 and R16000 systems. The directory structure on the originating system below CFDROOT would be:

```
$(CFDROOT)/
   LINUX32-GLIBC2.3/
      XEON/
         bin/
             Wind-US executable(s)
   LINUX64-GLIBC2.3/
      OPTERON/
         bin/
             Wind-US executable(s)
   SGI64-IRIX6.5/
      R14000/
         bin/
             Wind-US executable(s)
      R16000/
         bin/
             Wind-US executable(s)
   bin/
      Run scripts
   pvm/
      lib/
         LINUX32-GLIBC2.3/
             XEON/
                PVM executables
         LINUX64-GLIBC2.3/
             OPTERON/
                PVM executables
         SGI64-IRIX6.5/
             R14000/
                PVM executables
             R16000/
                PVM executables
```

When Wind-US is installed following the instructions in the *Wind-US Installation Guide*, the directory structure shown above is automatically created. Note that symbolic links may be used where appropriate. E.g., if only R14000 executables are available for SGI systems, the directory \$(CFDROOT)/SGI64-IRIX6.5/R16000 may be a symbolic link to \$(CFDROOT)/SGI64-IRIX6.5/R14000.

 $^{^{20}}$ CFDR00T is an environment variable set at login time, by running the cfd.login script in the user's .login file. For details see the instructions for installing the application or build distribution in the Wind-US Installation Guide.

9.5 Running Parallel Jobs

As noted earlier, running parallel jobs with Wind-US is remarkably simple. Assuming a parallel-capable executable is available for the system(s) being used, and the user's system and account have been set up as described above, the basic steps are:

- 1. Split the grid into multiple zones. Ideally, there should be one zone for each processor, and each zone should be the same size, assuming the processors have equal computing power. See Section 9.5.2 for additional details.
- 2. Create a multi-processing control (.mpc) file (see Section 9.6) listing the host names of the systems to be used.
- 3. Issue the wind command in one of the following forms.²¹
 - When running on a cluster or collection of distributed systems using rsh:

wind

• When running on a cluster or collection of distributed systems using ssh:

wind -usessh

• When running on a multi-processor system using PVM:

wind -mp

• When running on a multi-processor system using MPI:

wind -mp -mpmode MPI

For each of these, respond with "y" when prompted for whether or not you want to run in multi-processor mode. 22

More detail on various topics related to running parallel jobs with Wind-US is given in the following sections.

9.5.1 Command Line Options

The following wind script options are directly applicable to running Wind-US in parallel mode. See Section 8.1 for all the options.

-(no)parallel

Specifying -parallel indicates that the job is to be run in parallel mode, while -noparallel indicates serial mode. Parallel mode requires a multi-processing control (.mpc) file. If -parallel is specified, but an .mpc file doesn't exist, the user will be asked if serial mode should be used. If neither -parallel nor -noparallel is specified, and an .mpc file exists, the user will be asked if parallel mode should be used.

-mpmode mode

Message passing mode to be used when running in parallel: either PVM or MPI. Note that the pre-compiled executables available through IVMS were built without MPI message passing. To use MPI message passing, MPI must be pre-installed on your system(s) (unlike PVM, MPI is not distributed with Wind-US), and you'll need an executable that includes links to the MPI

 $^{^{21}}$ The commands shown here are the simplest forms. Additional wind script options, described in Section 8.1, may be used as needed.

²²The wording of this prompt is unfortunate. It really means "parallel mode", not necessarily on a single multi-processor system as defined earlier.

library. See the section "Installing the Build Distribution" in the Wind-US Installation Guide for instructions on creating the executable. The default message passing mode is PVM.

-(no)usessh When -usessh is specified, ssh/scp remote shell/copy commands will be

used when copying files between systems when running in parallel mode on a clusters or distributed systems. The default is to use rsh/rcp. For more details see the discussion of remote shell commands for parallel processing

in Section 9.3.

-(no)mp Specifying -mp indicates that a multi-processor machine (i.e., a single ma-

chine with multiple processors, like the CRAY or SGI Origin-2000) is being used, with either PVM or MPI message passing. The default is -nomp.

-nzones number Number of zones, used in MPI message passing mode. If not specified, the

Wind-US utility mpigetnzone will be used to get the number of zones from the common grid (.cgd) file. If mpigetnzone is not installed, the user will be

prompted for the number of zones.

9.5.2 Zone Size Considerations

Because synchronization takes place at the end of each cycle, total throughput is established by the processor that takes the longest to complete its assigned work. The optimum situation is to have all zones of equal size and have one processor for each zone. This gives maximum throughput and processor utilization, but is generally not achievable. If all zones cannot be close to the same size, a mixture of sizes is preferable. The case to avoid is a configuration with one zone of comparable size to the sum of the remaining zones. In this case, one can achieve at most a factor-of-two performance improvement regardless of the number of processors used. In general, if n is the number of points in the largest zone and N is the total number of points, the maximum possible speed up is N/n (assuming identical processors and similar algorithm specification).

Given a number of processors P with relative speeds p_i (larger p implies faster), and a number of zones N of sizes n_j , the assignment of work is done as follows.

- 1. Assign the largest zone j to processor 1 and compute $T_1 = n_j/p_1$.
- 2. Repeat step 1 for the remaining P-1 processors, assigning the largest remaining zone j to processor i and compute $T_i = n_i/p_i$.
- 3. If any zones remain to be assigned, locate processor i such that T_i is a minimum. Assign the largest remaining zone j to processor i, computing $T_i = T_i + n_j/p_i$.
- 4. Repeat step 3 for remaining unassigned zones.

Consider adding processors if T for any processor is significantly larger than the others, and that processor has more than one zone assigned.

The list output (.lis) file from a run will indicate what zones are assigned to what processor, and will have a report containing the utilization of each processor.

9.5.3 Checkpointing and Fault Tolerance

The flow (.cft) file contains the computed flow field results for all the zones, and is stored on the master. Since in parallel mode the zones are solved on separate processors, it's necessary to

periodically update the .cfl file on the master. By default, the frequency for doing this is once an hour (wall clock time), but this may be changed using the checkpoint directive in the .mpc file.

If a worker system fails due to either a system or network failure during the course of a run, the job will restart from the last checkpoint without the failed system. The automatic restart ability will be invoked as many times as necessary during a job until no more systems are available.

9.5.4 Intermediate Processing

At each checkpoint, the existing *.cfl* file is overwritten with the updated flow field. During long-running parallel jobs (or serial jobs, for that matter), it's sometimes desirable to do some intermediate processing, to examine how the solution is evolving, or to save snapshots of the results in an unsteady calculation.

The easiest way to do this is by using the SPAWN keyword, which allows user-specified processes to be run at user-specified intervals. Wind-US will temporarily stop while a spawned process is running, and continue when it finishes. One typical use of this capability is to run a user-written shell script that creates intermediate results from information in the .cft file. The description of the SPAWN keyword includes an example showing how to save intermediate .cft files for later post-processing.

By default, the .cfl file is automatically updated before starting the spawned process. This is in addition to the update of the .cfl file that's done at each checkpoint. Thus, if the SPAWN keyword is used, you may want to eliminate the normal checkpointing by specifying checkpoint none in the .mpc file.

9.5.5 Multiple Parallel Jobs

When running in parallel mode on a cluster or collection of distributed systems, the master system and all worker systems being used by a given user cannot be used by any other parallel job from the same user as long as the first job is active. A different user, however, can have a parallel job running simultaneously on the same systems, assuming that the memory, disk space, etc., are sufficient to support multiple jobs.

Note that in master mode the same originating system may be used to launch multiple parallel jobs, as long as the specified master and workers for each job don't overlap.

There are no restrictions on the number of parallel jobs for a given user on a multi-processor system (i.e., using the -mp option to the wind script; see Section 8.1), again assuming that the computer resources are available to support multiple jobs.

9.5.6 Stopping a Job

When a parallel Wind-US job finishes, the results files are updated on the master, various temporary files are removed on both the master and workers, and the run ends. If PVM message passing was used, PVM is stopped on the master and on all workers.

The methods for specifying when a parallel Wind-US run should stop are the same as for serial runs.

• The job will automatically end when the number of cycles specified by the user have been completed, or the solution converges.

- For non-interactive jobs, either a stop time or run time (depending on the queueing system being used) may be specified when the job is started using the *wind* script. See Section 8.1 for details.
- An NDSTOP file may be created in the Wind-US run directory to stop the job, as described in Section 7.9.

Because parallel jobs are often run during off-shift hours, using systems that are needed for other work during normal hours, scripts are supplied with Wind-US that may be executed by the Unix *cron* process to assure that jobs aren't inadvertantly run beyond a certain time. In the Wind-US distribution, these scripts are in the directory wind/bin/pvmkill. Four files are located there:

cronkill This file tells the continuous running job scheduler when to terminate processes. The first two digits on each line are the minute, the third digit is the hour, and following the *'s are the days when each of the commands will be executed (Monday = 1). The first command is the "nicest" way to kill the job, with the following two successively harsher. Note that this file must be edited so that output

goes to your directory and the paths for the scripts are correct.

pvmclean A script which terminates jobs in a relatively nice fashion.naskill A script which terminates jobs in a bit harsher fashion.

naspvmkill A script which terminates jobs in the meanest fashion.

To invoke these processes, copy the above scripts to each master you're using, edit cronkill appropriately, and insert these processes into the crontab on each master by entering²³

crontab cronkill

To check if this worked, enter

crontab -1

which will give a list of all your cron entries.

9.5.7 Multi-Processors vs Clusters and Distributed Systems

Experience has shown that the differences in the procedures for running on a multi-processor system (i.e., a single system with multiple CPUs), and on a cluster or collection of distributed systems, can be confusing. The following table is an attempt to summarize the differences.

	Multi-Processor	Cluster/Distributed Systems
Definition	Single system with multiple CPUs	Networked systems (with or without NFS-mounted home directories)
Message Passing	PVM or MPI	PVM
Required $wind$ Options	-mp, -mpmode MPI (if MPI is used)	None
Host List in $.mpc$ File	One host line, with $nproc > 1$	Multiple host lines
Multiple Jobs OK?	Yes	No

 $^{^{23}}$ Depending on how your system is configured, use of crontab may require root access.

9.6 Multi-Processing Control File

The multi-processing control file specifies the hosts that will be available as well as some miscellaneous options. If the Wind-US input data file name is *input.dat*, the name of the multi-processing control file must be *input.mpc*. When this file is present, the *wind* script will ask the user if they really want to use multi-processing mode.²⁴

Comments may be included in the file with the normal Wind-US comment indicator '/', or additionally '#'. Blank lines are ignored. Trailing comments are not allowed. The formats of the directives follow.

```
\verb|host {localhost | } name \} \ [\verb|nproc | n ]
```

host directives specify the names of the worker systems (given by the *name* parameter) that will be used to process zones. In general, there should be one host directive for each worker system. If a particular system appears more than once, each occurrence is treated as a unique system and will process assigned zones simultaneously. This is not advisable unless the system has multiple processors and sufficient memory.

As noted earlier, Wind-US uses the Unix hostname command to determine a system's name. Thus, in the host directive the specified name must be the same as the name returned by the hostname command on that system. For example, for a system with the fully-qualified name "workerbee.bigcompany.com", if hostname returns just the machine name "workerbee", then workerbee should be used for name in the .mpc file. If hostname returns the fully-qualified name "workerbee.bigcompany.com", then workerbee.bigcompany.com should be used for name.

The optional parameter $nproc\ n$ may be used to specify the number of processes to allow to run in parallel on the specified host. It is equivalent to repeating the host directive n times.

If no host entries appear in the multi-processing control file, the originating system will automatically be selected as the only host. When used on a system with sufficient memory and the assignment mode dedicated directive, the normal I/O associated with a single processor solution will be eliminated (except for checkpoints).

The special parameter localhost is used when running on a multi-processing system and the system name is unknown at the time of job submittal, such as for batch systems (like NQE) that can spawn to multiple systems or clusters of servers. Using localhost is preferred over not putting in any host directives because it assures that the scripts set up Wind-US consistently.

host entries should appear in the file in decreasing order of computational power. Wind-US assigns the most computationally intensive zones to the highest entries in the list.

The system that originates the job is not automatically included in the host list. If it is desired to also assign solution tasks to the originating system, it should have a host entry like any other system. For estimating purposes, the master process typically consumes less than one percent of the CPU time on the master host.

When rsh/rcp remote shell commands are used, the maximum number of hosts that may be specified is limited to 25, due to timeout issues that can occur with rsh/rcp. If more than 25 hosts are to be used, ssh/scp must be used for communication between the master and workers.

²⁴[As noted earlier, the wording of this prompt, and the terminology "multi-processing control file", is unfortunate. It really applies to all parallel jobs, not just those on a multi-processor system as defined earlier.

$\#master\ master_host\ [run_dir]$

This directive specifies the use of master mode, which allows a system different from the originating system to be used as the master.²⁵ The input parameter *master_host* specifies the name of the master system, and *run_dir* specifies the run directory to be used on the master. Like the system names in the **host** directive, the specified *master_host* must be the same as the name returned by the *hostname* command on that system.

If run_dir is not specified, the job will be run in the subdirectory logname, where logname is the user's login name, under a parent directory chosen from the following, in the order listed.

- PVM_TEMP, if the environment variable PVM_TEMP is defined
- /lscratch, if it exists
- /scratch*, if it exists, where * matches any string of 0 or more characters
- /data/local, if it exists
- /tmp, if it exists

When the job finishes, the output files (i.e., the .cfl file, .lis file, etc.) are copied back to the originating system, and everything is deleted from the run directory on the master, and the workers.

When master mode is used, the **-runinplace** wind script option is automatically set. Master mode may not be used in debug mode (i.e., with the **-debug** wind script option).

i/o {direct | <u>indirect</u>}

This directive specifies the type of I/O access that worker systems have to files on the master. The default is indirect, which means that workers do not have access to the files on the master, and that file I/O must therefore be done using message passing to/from the master process.

On multi-processor systems, however, i/o direct may be used to indicate that the worker processes may access the files directly, bypassing communication through the master process. This significantly reduces communication overhead and increases performance by as much as 10-40%.

There are a couple of things to be aware of when using the i/o direct option. First, it should only be used when running on a multi-processor system, not with clusters or distributed systems. And second, the maximum number of open files per process that is allowed (an operating system limit) must be large enough.

```
communication {direct | indirect}
```

The communication directive specifies how messages and boundary condition data are sent between worker systems. The default is direct, meaning that workers are able to communicate directly, without going through the master. Specifying indirect means that communication between workers must go through the master.

communication direct may not be used with assignment mode transient.

If the Wind-US executable was built using the -DF90 Fortran compiler option (necessary with compilers that don't support allocatable components in derived types), specifying communication direct has no effect; communication indirect is automatically used.

²⁵Note that the #master directive is an exception to the use of # as a comment indicator.

packmode {memory | memoryxdr | pointer | pointerxdr}

This directive specifies the packing mode used when transferring data between the master and workers.

memory The data to be transferred is sent just as it is stored in memory on the local

machine, and not XDR (External Data Representation) encoded. Thus, all the

systems must use the same internal data format.

memoryxdr This mode only applies to PVM message passing, and specifies that the data

being transferred is to be XDR encoded, allowing the systems to have different

internal data formats.

pointer The data being transferred is copied directly from memory, instead of being first

copied into a send buffer. During the packing process, the amount of data to be sent is determined, and pointers are used to identify the data itself. This is similar to the memory option, in that the data is not XDR encoded before being

sent, but should be faster.²⁶

pointerxdr This mode is currently the same as memoryxdr.

The default packing mode is memory for PVM message passing, and pointer for MPI message passing.

Note that when memory or pointer is used, since the data is not XDR encoded, the master and all workers must have the same internal data format. If a parallel job is being run on a collection of distributed systems with different internal data formats (which implies that PVM is being used, since MPI can only be used with Wind-US on multi-processor systems), the directive packmode memoryxdr must be specified in the .mpc file.

```
checkpoint {[every] {time minutes | count cycles} | none}
```

This directive specifies how often the worker systems transfer their flow field information to the flow file on the master system. In the event of a failure, the solution is automatically restarted from the last checkpoint. Specifying too small a number can result in very high network overhead and low throughput. A large number improves performance but can cause wasted time if a lot of network failures occur. If checkpoint none is specified, the flow field information is updated only at the end of the job. The default value is

```
checkpoint every 60 minutes
```

Note that if the SPAWN keyword is used in the input data (.dat) file, the flow field information is also updated before each spawned process, unless the NOCHECKPOINT option is specified.

```
assignment mode {dedicated | shared | transient | combined}
```

assignment mode controls how tasks are assigned to processors. There may be multiple appearances of this directive. Each one affects subsequent host entries up to the next assignment mode directive. A description of each mode follows.

²⁶This mode currently doesn't work on Linux systems with Intel compilers, due to a problem with character pointers and array temporaries.

dedicated Each task (zone) gets a unique Unix process on the target system. If a system must

process more than one zone, each will have a separate process, but only one will be allowed to run at a time unless multiple host entries are present for the system. This is the default mode and should not be changed unless there is insufficient

memory and swap space for the processes assigned to the host.

shared Unless a system must process more than one zone, this mode is the same as dedi-

cated. If more than one zone must be processed, only one Unix process is allocated and data for individual zones is swapped to and from local disk on the target system. This mode should be used only if the target system does not have sufficient

memory and swap space to contain the zones it needs to process.

transient This is similar to shared mode, in that when a system must process more than

one zone, only one Unix process is allocated. However, data for individual zones

is written back to the master processor instead of the local disk.

combined Like shared and transient modes, when a system must process more than one zone, only one Unix process is allocated. However, instead of writing data for

individual zones to the local disk or to the master, all zonal data is kept in memory.

task mode [dynamic | static]

When there are more tasks (i.e., zones) than processors, this directive may be used to predetermine which tasks are assigned to which processors.

The default procedure (i.e., when task mode is not used, or task mode dynamic is specified) is to start by ordering the tasks by the estimated amount of computational work each will require. Then, for n processors, the task requiring the most work is assigned to the processor specified by the first host directive in the .mpc file, the task requiring the second-most work is assigned to the second processor, etc., until the first n tasks have been assigned to the n processors. When a processor finishes its task, the next task in the queue (i.e. task n+1) is assigned to that processor. And so on, until all tasks have been assigned.

When task mode static is specified, the tasks are again ordered by the estimated work required. The first n tasks are assigned to the first n processors, just as above. But then, instead of waiting for a processor to become idle before assigning the next task, we continue assigning tasks, now starting with processor n and moving upwards in the list of processors. I.e., task n+1 is assigned to processor n, task n+2 is assigned to processor n-1, etc. This continues back and forth until all tasks have been assigned.

Note that if there are the same number of processors (or more) as there are tasks, this directive has no effect.

route {indirect | direct}

Controls how data is routed between the master and worker tasks when PVM message passing is used on a cluster or collection of distributed systems. This directive does not apply to MPI message passing, or to multi-processor systems.

indirect All data goes from the task on the local machine to the local PVM daemon, over the network to the remote PVM daemon, which forwards it to the remote task.

This is the standard data transfer procedure in PVM, uses UDP (User Datagram Protocol), and is scalable.

direct

All data goes directly from the local task to the remote task, bypassing the PVM daemons. This is implemented by setting the PVM option PvmRoute to PvmRoute-Direct, and uses TCP (Transmission Control Protocol) for transferring data. This takes more time to initially set up the TCP links, but is faster for subsequent data transfers.

It should be noted that this procedure is not scalable, and may fail if the number of zones is large. (Each TCP link requires a file descriptor, and the total number of file descriptors that is allowed is limited by the operating system.) However, if a direct link cannot be established, the indirect procedure through the PVM daemons will automatically be used.

#LOADLIMIT limit

The Wind-US/PVM initialization script will automatically eliminate workers that are deemed "too busy." A system is defined to be "too busy" when its 15 minute load factor, as reported by the Unix *uptime* command (the last number on the line) is greater than a certain limit (0.60 by default).²⁷

The load factor for each worker will be displayed in the list output file at the top with the other messages that occur during the preparation of the workers. The load factor will be displayed as a percentage (0.60 corresponds to 60%). Note that load factors in excess of 100% are possible. A message will also be displayed if the load factor exceeds the allowed threshold.

Occasionally, there is a problem with the *uptime* command and it reports a high load factor when there is no load on the system. To avoid this problem, the #LOADLIMIT directive²⁸ may be used to override the default value of 0.60. The parameter *limit* specifies the load limit for all hosts up to the next #LOADLIMIT directive. A #LOADLIMIT directive with no parameters restores the default load limit. This command should only be used when you know that including an overloaded host will not affect your job.

The following example illustrates the use of the #LOADLIMIT directive in the multi-processing control file.

```
# Next statement considers hosts ws1463 and ws1464 loaded
# only if their load factor exceeds 100%
#loadlimit 100
host ws1463
host ws1464
# The next statement restores the default load limit
#loadlimit
host ws1465
# Use a really high limit for ws1466 - disables the limit check
#loadlimit 9999
host ws1456
```

 $^{^{27}}$ Note that the load factor is checked only at initialization time. If the system becomes busy during a run . . . well . . . too bad.

²⁸Note that this directive is an exception to the use of # as a comment indicator.

Another way to modify the default load limit is to set the PVM_LOAD_LIMIT environment variable before you submit your job. For example, csh/tcsh users could do:

setenv PVM_LOAD_LIMIT 75

10 Keyword Reference

10.1 Text Conventions

In this manual, keywords are indicated by upper-case words in a fixed-pitch font, LIKE THIS. In the actual input file, however, they may be entered as upper or lower case. They may also be abbreviated, but Wind-US does *not* check for uniqueness. Therefore, specifying "A I" for arbitrary inflow is not a good idea. In cases where multiple options are available, the default (if one exists) is underlined.

User-specified input parameters are indicated by italics, *like this*. These user-specified parameters are usually numeric values, but may also be other keywords.

Most keywords consist of a single line, containing the keyword and its user-specified input parameters. Other keywords, like CHEMISTRY, indicate the start of a keyword block containing several lines, bracketed by starting and ending keywords. Within a keyword block, only individual keywords relevant to that block may be used. Keyword blocks are indicated by the word "block" in parentheses. Some keywords have several options, making them long and complex. These are sometimes split into two (or more) lines for display purposes, with a "\" at the end of the line being continued, but must nevertheless be on a single line in the input data file.

The various keywords and keyword blocks are listed alphabetically, with each one starting a new page. The overall syntax for each keyword or keyword block is shown in a box at the top of the page, with the details following.

In addition, the following documentation conventions are used:

- The "or" symbol; used to separate multiple choices
- Delimiters surrounding optional entry(s)
- {} Delimiters surrounding multiple entries when exactly one of them is required

10.2 Zone Selection

Many of Wind-US's capabilities may be specified on a zone-by-zone basis. Keywords used to enable these capabilities may include a zone selector at the end of the keyword command in the input data file. The keywords for which this type of specification is valid include a zone_selector format specifier in their description. The zone_selector must be of the following form:

[ZONE] zone-list

where zone-list is of the form:

range1[, range2[, ... rangen]]

A range is of the form:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

begzone: Selects all zones from begzone to MAXZONE, the total number of zones in the

grid file

: endzone Selects all zones from 1 to endzone

ALL Selects all zones

For some keywords that turn on capabilities that may be selected by zone, the *zone_selector* is optional (indicated by the syntax [*zone_selector*]). In this case, omitting the zone specification defaults to using the capability in all zones. You should therefore change the default selection before changing individual zones, as changing the default will reset any zones which have been individually set earlier in the input file.

Example

To turn off (i.e., perform no iterations) in zones 5, 6, and 7, but still pass the information in these zones to the adjacent zones, and to use 10 iterations per cycle in all other zones, use the following sequence of keywords:

```
ITERATIONS PER CYCLE 10
ITERATIONS PER CYCLE -1 ZONE 5:7
```

Note that reversing the order of the keywords would not work because resetting the default will override the initial selection.

10.3 Keyword Details

The following is a comprehensive, alphabetical list of all keywords that are valid in the Wind-US input data file. Every attempt was made to build keyword inputs from intuitive, English-language words, and to adhere to some general rules of construction for the keyword commands in the data file.

Many of the keywords apply to either structured grids (e.g., ACTUATOR) or unstructured grids (e.g., Q LIMIT), but not both. For these keywords, the type of grid that the keyword applies to is listed above the box showing the keyword syntax.

Several other keywords apply to both types of grids, but with different options or syntaxes (e.g, IMPLICIT). For these keywords, typically two syntax boxes are shown, again with labels specifying the grid type. In general, the detailed description of these keywords is divided into separate sections for structured and unstructured grids.

The remaining keywords apply to both types of grids. If the keyword syntax box isn't labeled with the words "Structured Grids" or "Unstructured Grids" (e.g., FREESTREAM), that keyword applies to both types of grids, with (mostly) the same options and syntax. Some specific options may be limited to structured or unstructured grids, and these are noted in the text.

ACCELERATE — Convergence acceleration (block)

Structured Grids

```
ACCELERATE ZONE n [CFLRAMP START iter_1 END iter_2] [SECOND START iter_1 END iter_2] [FOURTH START iter_1 END iter_2] ENDACCELERATE
```

Unstructured Grids

```
ACCELERATE ZONE n [CFLRAMP START iter_1 END iter_2] ENDACCELERATE
```

The objective of Wind-US's convergence acceleration scheme is to approach the steady state solution more quickly by allowing the use of a large CFL number early in the calculation.

Structured Grids

The ZONE keyword identifies the zone within which convergence acceleration is to be used. If convergence acceleration is to be used in multiple zones, and/or if different starting and ending iteration values are to be used in different zones, multiple ZONE keywords must be specified.

The CFL number will be ramped down linearly from its starting value, specified using the CFL# keyword, to 1.0. The ACCELERATE keyword does not work when CFL# MODE 3 (i.e., the time step calculation procedure from OVERFLOW) is used.

With structured grids, first- and second-order smoothing are normally applied to maintain numerical stability. The amount of smoothing, when it is applied, and how it varies with iteration is controlled by the SMOOTHING and/or BOUNDARY-DAMP keywords, the $iter_1$ and $iter_2$ values specified with ACCELERATE, and the mode used with TEST option 49.

For the second-order smoothing,

mode Second-Order Smoothing ≤ 2 Smoothing varies with ite

< 2 Smoothing varies with iteration i, as follows:

```
i \leq iter_1 Smoothing applied as determined by the SMOOTHING keyword iter_1 < i < iter_2 Smoothing coefficient decreased linearly to 0 No smoothing applied
```

2 Smoothing varies with iteration i, as follows:

```
\begin{array}{ll} i \leq iter_1 & \text{Smoothing applied as determined by the $\tt SMOOTHING} \text{ and} \\ & \texttt{BOUNDARY-DAMP} \text{ keywords} \\ iter_1 < i < iter_2 & \text{Smoothing coefficient decreased linearly to 0} \\ i \geq iter_2 & \text{No smoothing applied} \end{array}
```

3 No smoothing applied

And for the fourth-order smoothing,

```
Second-Order Smoothing
mode
 < 2
       Smoothing applied as determined by the SMOOTHING keyword
 2
       Smoothing varies with iteration i, as follows:
           i < iter_1
                                          No smoothing applied
           iter_1 < i < (iter_1 + iter_2)/2
                                         Smoothing coefficient increased linearly from 0
                                          to value determined by the SMOOTHING and/or
                                          BOUNDARY-DAMP keywords
           (iter_1 + iter_2)/2 < i < iter_2
                                          Smoothing coefficient decreased linearly to 0
           i \geq iter_2
                                          No smoothing applied
```

3 Smoothing applied as determined by the ${\tt SMOOTHING}$ and ${\tt BOUNDARY-DAMP}$ keywords

If $iter_1$ and $iter_2$ aren't specified, the following default values are used.

$\underline{\text{Option}}$	$\underline{iter_1}$	$iter_2$
SECOND	1	150
FOURTH	50	500
CFLRAMP	50	500

Recommended values for the second- and fourth-order smoothing parameters val_1 and val_2 specified with the SMOOTHING keyword, and for the starting CFL number, are:

Dimensions	$\underline{val_1}$	$\underline{val_2}$	CFL#
2D	0.1	0.03	10 – 15
3D	0.06	0.01	2.5

<u>Example</u>

In the following example, convergence acceleration is applied in zones 3 and 4. Second-order smoothing is applied between iterations 1 and 200, and fourth-order smoothing is applied between the default values of iterations 50 and 500. The CFL number specified with the CFL# keyword will be used for the first 250 iterations, then ramped down to a value of 1.0 between iterations 250 and 500.

```
SMOOTHING SECOND 0.06 FOURTH 0.01 SMLIMT 0.
ACCELERATE
ZONE 3
ZONE 4
SECOND START 1 END 200
CFLRAMP START 250 END 500
ENDACCELERATE
TEST 49 2
```

See Also: SMOOTHING, BOUNDARY-DAMP, CFL#, TEST 49

Unstructured Grids

The ZONE keyword identifies the zone within which convergence acceleration is to be used. If convergence acceleration is to be used in multiple zones, and/or if different starting and ending iteration values are to be used in different zones, multiple ZONE keywords must be specified.

 $iter_1$ and $iter_2$ specify starting and ending iteration values for varying the CFL number. Just as for structured grids, for unstructured grids the CFL number will be ramped down linearly from its starting value, specified using the CFL# keyword, to 1.0. If the CFLRAMP keyword isn't specified, default values of 50 and 500 are used, for $iter_1$ and $iter_2$.

For unstructured grids, however, no variation in smoothing is done.

See Also: CFL#

ACTUATOR | SCREEN — Actuator disk / screen model (block)

Structured Grids

```
{ACTUATOR | SCREEN}
   ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \
        [SUBSET I range J range K range]
   ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \
        [SUBSET I range J range K range]
  TURNING {CONSERVE {ANGLE | PARALLELU} | \
            ZERO PARALLELU | \
            {SOLIDBODY | VORTEX} val \ x_c \ y_c \ z_c | \
            SPECIFY ANGLE \alpha [ROTATE \beta]}
  TIP-EFFECT r1 r2 r3 r4
  POWER {{DPS | DPT | DPOWER} val | \
          TURNING | \
          {SOLIDBODY | VORTEX} val x_c y_c z_c}
  EFFICIENCY {ETA val | \
                CLOSS val | \
                VORTEX val | \
                SCREEN {NORMAL | TOTAL} SOLIDITY sol}
{ENDACTUATOR | ENDSCREEN}
```

This keyword enables the user to model an actuator disk or screen by specifying a discontinuous change in properties across a zone boundary or portion of a zone boundary. The following restrictions apply:

- This keyword applies to structured grids only.
- The BOUNDARY TVD FACTOR 0 keyword option should be used for all actuator disk and screen boundaries.
- Screens require zero work (POWER DPOWER 0.)
- Only one storage location for center of rotation, the last one encountered is used for all centers.

The various elements of the ACTUATOR | SCREEN input block are defined as follows:

```
{ACTUATOR | SCREEN}
```

Defines the beginning of the actuator or screen block.

```
ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \ [SUBSET I range J range K range]
ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \ [SUBSET I range J range K range]
```

These two lines define the location of the actuator disk or screen. The relevant zones are given by the values of iz1 and iz2, and the relevant boundaries within zones iz1 and iz2 are specified via the BOUNDARY keyword parameter.

iz1 Zone to which increments will be added when passing information to iz2

iz2 Zone receiving positive increments, increments will be subtracted when passing information back to zone iz1

The SUBSET parameter may be used to specify that the change in properties occurs only over a part of the zone boundary. Otherwise, it is assumed that the change occurs over the entire boundary. The *range* parameters define the part of the zone boundary over which the change occurs, and take one of the following forms:

index1 index2 Starting and ending indices in the specified direction. LAST may be used for the last index.

ALL Equivalent to 1 LAST.

The starting and ending indices for the appropriate I, J, or K parameter (depending on the boundary specified) must be the same, and correspond to that boundary. In addition, for two-dimensional cases, the K parameter must be specified as either K 1 1 or K ALL.

```
TURNING {CONSERVE {ANGLE | PARALLELU} | \ ZERO PARALLELU | \ {SOLIDBODY | VORTEX} val \ x_c \ y_c \ z_c | \ SPECIFY ANGLE \alpha [ROTATE \beta]}
```

Defines the net change in parallel velocity across the zone boundary.

CONSERVE Conserves the net flow angle (ANGLE) or the parallel velocity components

 $({\tt PARALLELU})$ across the zone boundary. (The ${\tt ANGLE}$ option is currently not

implemented.)

ZERO PARALLELU Sets the parallel components of velocity across the zone boundary to zero

SOLIDBODY Defines a solidbody rotation increment to the parallel velocity, where:

val Rotation rate in degrees/second (positive by right hand rule).

(Note: prior to Wind-US 2.184 this was in radians per second. Input data files using this keyword with earlier versions of Wind-US will need to be changed if used with Wind-US

2.184 or later.)

 x_c, y_c, z_c Center of rotation (must be in the plane, requires the zone boundary to lie in a constant x, y, or z plane) (inches)

VORTEX Defines free vortex flow increment to parallel velocity, where:

val Vortex strength κ (ft²/sec), where $\kappa = \omega a^2 = \Gamma/2\pi$, and Γ is the circulation, ω is the rotation rate of the solidbody core, and a is the radius of the solidbody core (required to avoid P = 0 at axis), $a^2 = \kappa^2 \rho/(0.9P_{\infty})$ (assumes $P_{min} = 0.1P_{\infty}$)

 x_c, y_c, z_c Center of rotation (inches)

SPECIFY ANGLE Allows the user to specify the flow turning angle.

alpha The flow angle giving the rotation of the iz2 boundary normal, projected onto the x-y plane, about the z-axis (degrees)

beta An optional rotation of the resulting vector about the y-axis

TIP-EFFECT r1 r2 r3 r4

Forces increments to go to zero at hub and/or tip to avoid solution discontinuities at the boundaries. A scalar, (0-1) multiplies the turning and power when this option is on. This is required for engine face models (where the wall velocity at the tip must be zero in the diffuser frame of reference). r1-r4 define linear regions ranging from 0 to 1 between r1 and r2, and from 1 to 0 between r3 and r4. r1, r2, r3, and r4 define the distance from the center of rotation (inches).

This keyword requires that TURNING SOLIDBODY, TURNING VORTEX, or POWER SOLIDBODY be specified.

POWER {{DPS | DPT | DPOWER} val | \ TURNING | \ {SOLIDBODY | VORTEX} val x_c y_c z_c }

Defines the power increment across the zone boundary. Screens require setting the power to zero. i.e., $POWER\ DPOWER\ 0$.

DPS val specifies the static pressure increment across the actuator boundary

(psi). Requires that the efficiency be specified, using EFFICIENCY ETA.

DPT val specifies the total pressure increment across the actuator boundary (psi). Requires that the efficiency be specified, using EFFICIENCY ETA.

DPOWER val specifies a (constant) power per unit area increment. (ft-lb/sec-ft²). (Corresponds to unsteady (rotor) free vortex turning)

$$\mathtt{DPOWER} = \rho u c_p (T_{t2} - T_{t1})$$

TURNING

Specifies work corresponding to the net turn across the zone boundary (specified in the TURNING element). Assumes all turning is done in an unsteady process (by the rotor), i.e., no stator.

$$dW = c_n(T_{t2} - T_{t1}) = \omega r(w_2 - w_1)$$

where ω is the rotation rate of the rotor, r is the local radius from the center of the rotor, and w is the local circumferential velocity.

This option requires that TURNING SOLIDBODY be specified.

SOLIDBODY | VORTEX

Defined as in the TURNING element. This defines the turning accomplished by the rotor. The net turning may be altered by another process (e.g., by a stator).

$$dW = c_p(T_{t2} - T_{t1}) = \omega r(w_2 - w_1)$$

Note: Currently vortex turning (i.e., POWER VORTEX) is not allowed. This would correspond to constant work across the rotor. However, currently, the procedure used to eliminate the vacuum at the core (setting $P_{min}=0.1P_{\infty}$) makes the work input independent of the strength of the vortex, so the user could not vary the work input by changing κ .

Defines the efficiency of the actuator disk or screen.

ETA Compressor efficiency, $val = [(P_{t2}/P_{t1})^{(\gamma-1)/\gamma} - 1]/[(T_{t2}/T_{t1}) - 1]$

CLOSS Loss coefficient, $val = (P_{t1} - P_{t2})/q$, where $q = \rho U^2/2$ (*U* based on normal Mach number)

VORTEX Free vortex total pressure loss. The value val is the maximum value of $(P_{t2}-P_{t1})/P_{tinf}$ (i.e., the loss at the center of the vortex). A linear distribution is assumed from the vortex center to the radius a, where a is determined by the strength value specified using TURNING VORTEX, or directly using TEST 180.

This option requires that TURNING VORTEX be specified.

SCREEN Use screen loss relations to define total pressure loss, where

NORMAL Use normal component of Mach number

TOTAL Use total Mach number. (This option is not currently implemented.)

Solidity of screen = $A_b/(A_b + A_o)$, where A_b is the blocked area, and A_o is the open area.

If solidity is specified, the screen loss coefficient associated with the screen model is defined by the solidity correlation of Cornell (1958), unless the optional CLOSS value is specified.

The screen model is not intended for use with choked screens, where the screen is significantly limiting the mass flow rate. During the solution start-up phase, it may be necessary to specify a low solidity, then increase it to the desired value to avoid strong choking in transients.

This option requires that the power be zero. i.e., POWER DPOWER 0.

```
{ENDACTUATOR | ENDSCREEN}
```

Ends actuator or screen input block

Examples

The following examples illustrate the use of the ACTUATOR | SCREEN input block for an engine face and for a screen.

Engine face model

```
ACTUATOR

ZONE 1 BOUNDARY IMAX

ZONE 2 BOUNDARY I1

TURNING SOLIDBODY 240000. 312. 54. 0.

TIP-EFFECT 5. 5.1 39.8 40.0

POWER TURNING

EFFICIENCY ETA 0.85

ENDACTUATOR
```

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```
BOUNDARY TVD FACTOR O ZONE 1 BOUNDARY IMAX BOUNDARY TVD FACTOR O ZONE 2 BOUNDARY I1

Screen

SCREEN

ZONE 3 BOUNDARY K1

ZONE 2 BOUNDARY IMAX

TURNING ZERO PARALLELU

POWER DPOWER 0.0

EFFICIENCY SCREEN NORMAL SOLIDITY 0.1

ENDSCREEN
```

BOUNDARY TVD FACTOR O ZONE 3 BOUNDARY K1
BOUNDARY TVD FACTOR O ZONE 2 BOUNDARY IMAX

See Also: BOUNDARY TVD, TEST 180

ARBITRARY INFLOW — Arbitrary inflow (block)

Structured Grids

```
{ARBITRARY INFLOW | DIFFUSER INFLOW}
    [STATIC | TOTAL]
    [HOLD_TOTALS | HOLD_CHARACTERISTICS]
    [DIRECTION {SPECIFIED | CYLINDRICAL [X|Y|Z] | NORMAL [TO INFLOW PLANE] | \
                  ALONG [GRID LINES] }]
    [UNIFORM [MPT \alpha \beta [val\_k [val\_om]]]
       [sp_1 \ sp_2 \ \ldots sp_n]
    [IJK_RANGE [FROZEN] imin\ imax\ jmin\ jmax\ kmin\ kmax\ M\ P\ T\ \alpha\ \beta\ [val_k\ [val_om]]
       [sp_1 \ sp_2 \ \ldots sp_n]
    [UNSTEADY var_name freq ampl phase]
    [XYZ_RANGE [BLEND {XMIN | XMAX}] xmin xmax ymin ymax zmin zmax \
                                             MPT \alpha \beta [val\_k [val\_om]]
       [sp_1 \ sp_2 \ \ldots sp_n]
    [RTZ_RANGE [BLEND {ZMIN | ZMAX}] rmin rmax tmin tmax zmin zmax \
                                             MPT \alpha \beta [val\_k [val\_om]]
       [sp_1 \ sp_2 \ \dots sp_n]
    [TURBULENT [INFLOW] [MAGNITUDE] vpmag {FPS|MEAN|PERCENT} [SIZE nsiz] \
       [SEED seed] [PEAK_K kpeak [PER_FOOT]] [BL_HEIGHT blhgt]]
    [{VORTEX | SOLIDBODY | ROTATESOLID} M_n P T \alpha \beta xc yc zc \backslash
       \{dw1 \mid dw1 \ dw2 \ dw3\}
    [USERSPEC fs bl1 bl2 npts
       y1 MPT \alpha \beta
       y2~M~P~T~\alpha~\beta
       yn M P T \alpha \beta
    [USERCHEM fs bl1 bl2 npts
       y1 MPT \alpha \beta
           sp_1 \ sp_2 \ \dots sp_n
       y2 MPT \alpha \beta
           sp_1  sp_2  \dots sp_n
       yn MPT \alpha \beta
           sp_1 \ sp_2 \ \dots sp_n
    [USERKE fs\ bl1\ bl2\ npts
       y1 MPT \alpha \beta val_k [val_om]
       y2 MPT \alpha \beta val_k [val_om]
       yn \ M \ P \ T \ \alpha \ \beta \ val\_k \ [val\_om]]
    [USERCHEMKE fs bl1 bl2 npts
       y1 MPT \alpha \beta
           sp_1  sp_2  \dots sp_n  val_k  [val\_om]
       y2 MPT \alpha \beta
           sp_1  sp_2  \dots sp_n  val_k  [val\_om]
       yn M P T \alpha \beta
           sp_1  sp_2  \dots sp_n  val\_k  [val\_om]]
[ENDINFLOW]
```

Unstructured Grids

```
{ARBITRARY INFLOW | DIFFUSER INFLOW}
   [STATIC | TOTAL]
   [HOLD_TOTALS]
   [DIRECTION {SPECIFIED | CYLINDRICAL [X|Y|Z] | NORMAL [TO INFLOW PLANE]}]
   ZONE n
   [UNIFORM [MPT\alpha\beta]]
       [sp_1 \ sp_2 \ \ldots sp_n]
   [USURFACE [FROZEN] surface M P T \alpha \beta [sp_1 \ sp_2 \ ... sp_n]]
   [UNSTEADY var_name freq ampl phase]
   [XYZ_RANGE [BLEND {XMIN | XMAX}] xmin xmax ymin ymax zmin zmax \
                                          MPT \alpha \beta [val_k [val_om]]
      [sp_1 \ sp_2 \ \dots sp_n]
   [RTZ_RANGE [BLEND {ZMIN | ZMAX}] rmin rmax tmin tmax zmin zmax \
                                          MPT \alpha \beta [val\_k [val\_om]]
       [sp_1 \ sp_2 \ \ldots sp_n]
[ENDINFLOW]
```

Several options are available to set boundary conditions at arbitrary inflow boundaries. The default is uniform inflow (i.e., no boundary layer) at the conditions that are set using the FREESTREAM keyword. Other options are selected by using either of the equivalent keywords ARBITRARY INFLOW or DIFFUSER INFLOW.

The remaining lines select the specific type of inflow data to be provided. Discussion of the input data for each of these options is grouped into sections labeled "Control Functions", "Condition Specification", and "Special Capabilities". These keywords can start in any column. Generally, they should be indented from the ARBITRARY INFLOW keyword to set them apart.

This keyword may also be used to initialize (or reinitialize) the flow conditions within the specified zone, as described in Section 3.6.

Note: The GAS keyword and the CHEMISTRY keyword block, if used, must come before the ARBITRARY INFLOW keyword block in the input data (.dat) file.

Structured Grids

Control Functions

ENDINFLOW

This optional keyword may be used to end the arbitrary inflow definition.

```
STATIC | TOTAL
```

Arbitrary inflow conditions specified after this keyword will be considered as static or total, depending which of these is set. By default, the flag is set to the value from the FREESTREAM keyword.

When TOTAL is specified, the static conditions are computed using formulas for perfect gases. If total conditions must be specified for real gas flows, the GAS keyword should be used to specify a value for the specific heat ratio γ that is consistent with the real gas mixture being used.

```
HOLD_TOTALS | HOLD_CHARACTERISTICS
```

HOLD_TOTALS indicates that total temperature and local flow angles are to be held at their specified values. HOLD_CHARACTERISTICS indicates that characteristic values are to be held constant. The option specified will be applied at all the arbitrary inflow regions in the zone, and will remain in effect for all following ZONE keywords.

The default for structured grids is to hold characteristic values constant.

These keywords only work in conjunction with the UNIFORM, IJK_RANGE, VORTEX, SOLIDBODY, and ROTATESOLID keywords.

The total pressure is always held fixed, whether HOLD_TOTALS is specified or not. Although the Mach number is specified with the UNIFORM and IJK_RANGE keyword parameters, it may be adjusted during the boundary condition treatment.

Note that the HOLD_TOTALS keyword in the ARBITRARY INFLOW keyword block applies to arbitrary inflow boundaries only. See the HOLD keyword for information on holding conditions at freestream boundaries with inflow.

Note also that the syntax is slightly different for arbitrary inflow and freestream boundaries. For arbitrary inflow boundaries, HOLD_TOTALS and HOLD_CHARACTERISTICS are used in the ARBITRARY INFLOW keyword block, with an underscore. For freestream boundaries, HOLD TOTALS and HOLD CHARACTERISTICS are used, without an underscore.

```
DIRECTION {SPECIFIED | CYLINDRICAL [X|Y|Z] | NORMAL [TO INFLOW PLANE] | \
           ALONG [GRID LINES]}
```

The DIRECTION keyword indicates how the flow angle is to be set at an inflow plane. The options for setting the flow direction are:

SPECIFIED Set the flow at the angles of attack and yaw specified elsewhere in the

ARBITRARY INFLOW block. The angles are defined in the same way as for the FREESTREAM keyword. This is the default.

CYLINDRICAL [X|Y|Z] This is the same as SPECIFIED, except that the angles are used to specify velocities in cylindrical coordinate directions. I.e., if V is the inflow

> velocity magnitude, α and β are related to the radial, circumferential, and axial cylindrical velocity components by:

> > $v_r = V \sin \beta$

 $v_{\theta} = V \cos \beta \sin \alpha$ $v_z = V \cos \beta \cos \alpha$

The cylindrical axis of rotation must correspond to one of the three Cartesian axis directions, as specified by the option X, Y, or Z. The default is X.

Set the flow normal to the inflow plane.

NORMAL

ALONG

Set the flow in the direction of the grid lines intersecting the inflow plane.

Angles specified with the DIRECTION keyword will override any angles of attack or yaw specified when setting boundary condition values with the UNIFORM, IJK_RANGE, VORTEX, SOLIDBODY, and ROTATESOLID keywords. However, if UNIFORM is used without specifying the flow conditions, the angles specified with the FREESTREAM keyword will be used, and the DIRECTION keyword will have no effect. The DIRECTION keyword also does not affect flow angles in profiles specified with the USERSPEC, USERCHEM, USERKE, or USERCHEMKE keywords.

The DIRECTION keyword will not modify the flow angles that are set when ARBITRARY INFLOW is being used to initialize (or reinitialize) flow conditions within a zone.

The DIRECTION option used will remain in effect for all following ZONE keywords.

```
{\tt ZONE}\ n
```

This keyword, which *must* be specified, identifies the zone for which inflow conditions are being set. For example, if zone 2 is an internal jet, conditions other than freestream may be desired at the inflow to zone 2.

Within the ARBITRARY INFLOW keyword block, the ZONE keyword must come before any "Condition Specification" keyword for that zone, but after any HOLD_TOTALS, HOLD_CHARACTERISTICS, or DIRECTION keyword for that zone. In addition, inflow conditions may only be specified for one zone at a time.

Example

The following ARBITRARY INFLOW block specifies that total conditions are to be held constant at arbitrary inflow surfaces in zones 1 and 2, with M=3.5, $P_T=251.15$ psi, and $T_T=1167.9$ °R. In zone 3, characteristic values are to be held constant at arbitrary inflow surfaces, consistent with the flow conditions given with the FREESTREAM keyword.

```
ARBITRARY INFLOW

TOTAL

HOLD_TOTALS

ZONE 1

UNIFORM 2.5 251.15 1167.9 0.0 0.0

ZONE 2

UNIFORM 2.5 251.15 1167.9 0.0 0.0

HOLD_CHARACTERISTICS

ZONE 3

UNIFORM

ENDINFLOW
```

Condition Specification

```
UNIFORM [M\ P\ T\ lpha\ eta [val\_k [val\_om]]] [sp_1\ sp_2\ \dots sp_n]
```

This keyword is used to specify uniform flow at arbitrary inflow boundaries, at the flow conditions listed below. If the flow conditions are omitted, those specified with the FREESTREAM keyword are used.

M

Mach number

Pressure, psi $T \qquad \qquad \text{Temperature, } ^\circ \mathbf{R} \\ \alpha, \, \beta \qquad \qquad \text{Angles of attack and yaw, relative to the Cartesian } x \text{ direction, in degrees} \\ sp_1, sp_2, \ldots, sp_n \qquad \text{Species mass fractions. These are required for real gas flows. The order for these values must be consistent with the order in which the species are listed with the SPECIES keyword in the CHEMISTRY keyword block.}$

Pressure and temperature are static or total, depending on whether STATIC or TOTAL is specified.

For structured grids, when the SST turbulence model is being used (see the TURBULENCE keyword), val_k and val_om may be used to specify inflow turbulence levels. You may specify either val_k , or val_k and val_om , but not val_om by itself. Note that if these values are being specified, the TURBULENCE keyword must come before the ARBITRARY INFLOW keyword block in the input data (.dat) file.

The following options are possible:

 $val_{-}k > 0$ The turbulent kinetic energy k and the specific dissipation rate ω are specified directly, with

$$k = val_k \text{ (ft}^2/\text{sec}^2)$$

 $\omega = val_om \text{ (1/sec)}$

The turbulent viscosity ν_t is then equal to k/ω .

 $val_k < 0$ The turbulence intensity is set equal to $|val_k|$, expressed as a percentage of the inflow velocity U, where U is computed from the specified values of M and T. Thus, the turbulent kinetic energy is computed as

$$k = 1.5(0.01 | val_k | U)^2$$

The turbulent viscosity ν_t is automatically set equal to $0.001\nu_l$, where ν_l is the laminar viscosity, and the specific dissipation rate is computed as $\omega = k/\nu_t$.

 $val_om < 0$ The specific dissipation rate ω is set equal to val_om percent of U/L_{ref} , where U is computed from the specified values of M and T, and L_{ref} is the reference length from the grid (.cgd) file. Thus

$$\omega = 0.01 \ |val_om| \ \frac{U}{L_{ref}}$$

The turbulent viscosity ν_t is set to the same percentage of the laminar viscosity.

$$\nu_t = 0.01 |val_om| \nu_l$$

The turbulent kinetic energy is then computed as $k = \omega \nu_t$.

If inflow turbulence levels are not specified using one of the above options, or if $val_k = 0$, default values are computed from

$$\omega = 10U/L_{ref}$$

$$\nu_t = 0.001\nu_l$$

$$k = \omega\nu_t$$

Note that

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- If val k > 0, a positive value must be specified for val om.
- If $val_k \leq 0$, val_om should not be specified.
- If $val_om < 0$, a value must also be specified for val_k , but it is ignored.

Example

```
ARBITRARY INFLOW
UNIFORM 1.1 100. 900. 10. 0.
ENDINFLOW
```

```
IJK_RANGE [FROZEN] imin\ imax\ jmin\ jmax\ kmin\ kmax\ M\ P\ T\ \alpha\ \beta [val_k [val_om]] [sp_1\ sp_2\ \dots sp_n]
```

For structured grids, this keyword allows specification of inflow conditions over an arbitrary range of i, j, and k indices on any computational boundary plane. The user specifies the minimum and maximum i, j, and k indices which describe the region, followed by the flow conditions to be applied, as follows:

imin, imax	Minimum and maximum i indices bounding the region
jmin, jmax	Minimum and maximum j indices bounding the region
kmin, kmax	Minimum and maximum k indices bounding the region
M	Mach number
P	Pressure, psi
T	Temperature, ${}^{\circ}R$
α,eta	Angles of attack and yaw, relative to the Cartesian x direction, in degrees
sp_1, sp_2, \ldots, sp_n	Species mass fractions. These are required for real gas flows. The order for these values must be consistent with the order in which the species are listed with the SPECIES keyword in the CHEMISTRY keyword block.

Pressure and temperature are static or total, depending on whether STATIC or TOTAL is specified. There are no defaults for the index ranges.

The FROZEN option may be specified to freeze the inflow conditions over the indicated index range at their current values. Note that for a restart case (i.e., when a .cfl file already exists), the "current values" are those in the .cfl file, not those specified with the IJK_RANGE keyword. For an initial run (i.e., when a .cfl file does not exist), the flow conditions will be frozen at the conditions specified with the IJK_RANGE keyword.

If the SST turbulence model is being used (see the TURBULENCE keyword), val_k and val_om may be used to specify inflow turbulence levels. The various options are described above under the UNIFORM keyword.

A combination of up to 4000 IJK_RANGE, XYZ_RANGE, and RTZ_RANGE regions and (for unstructured grids) USURFACE surfaces may be specified. This is useful when specifying a boundary layer profile at an inflow boundary, or along solid walls during the flowfield initialization process, as described in Section 3.6 starting on p. 39.

UNSTEADY var_name freq ampl phase

This keyword allows the user to specify unsteady arbitrary inflow conditions. It must be used with (and follow) the IJK_RANGE keyword. Up to ten different perturbations to the inflow conditions may be specified and will be superimposed to create unsteadiness centered about the conditions given with the IJK_RANGE keyword.

var_name One of the keywords MACH, PRESSURE, TEMPERATURE, ALPHA, BETA, or VELOCITY

freq Frequency of the perturbation in Hertz

ampl Amplitude of the perturbation in appropriate variable units

phase Phase angle of the perturbation in degrees

UNSTEADY may only be used when a constant time step is specified with the CFL# keyword, and only for a perfect gas.

Note that you may specify multiple, independent pairs of IJK_RANGE and UNSTEADY keywords.

```
XYZ_RANGE [BLEND {XMIN | XMAX}] xmin\ xmax\ ymin\ ymax\ zmin\ zmax \setminus M\ P\ T\ \alpha\ \beta\ [val\_k\ [val\_om]] [sp_1\ sp_2\ \dots sp_n]
```

This keyword is intended for use during the flowfield initialization process (Section 3.6), and allows specification of initial conditions in a specified Cartesian bounding box. The user specifies the bounding box coordinates, followed by the flow conditions to be applied, as follows:

xmin, xmax	x_{min} and x_{max} coordinates of bounding box
ymin, ymax	y_{min} and y_{max} coordinates of bounding box
zmin, zmax	z_{min} and z_{max} coordinates of bounding box
M	Mach number
P	Pressure, psi
T	Temperature, ${}^{\circ}R$
α,eta	Angles of attack and yaw, relative to the Cartesian x direction, in degrees
sp_1, sp_2, \ldots, sp_n	Species mass fractions. These are required for real gas flows. The order for these values must be consistent with the order in which the species are listed with the SPECIES keyword in the CHEMISTRY keyword block.

Pressure and temperature are static or total, depending on whether STATIC or TOTAL is specified. There are no defaults for the bounding box coordinates.

If BLEND XMIN or BLEND XMAX is specified, the initial conditions will smoothly transition from the specified values at x_{min} to freestream values at x_{max} , or from the specified values at x_{max} to freestream values at x_{min} , respectively. If BLEND is not specified, the initial conditions will be uniform, at the specified values. This option is only available for perfect gas flows.

If the SST turbulence model is being used (see the TURBULENCE keyword), val_k and val_om may be used to specify inflow turbulence levels. The various options are described above under the UNIFORM keyword.

A combination of up to 4000 IJK_RANGE, XYZ_RANGE, and RTZ_RANGE regions and (for unstructured grids) USURFACE surfaces may be specified. This is useful when specifying a boundary layer

profile at an inflow boundary, or along solid walls during the flowfield initialization process, as described in Section 3.6 starting on p. 39.

```
RTZ_RANGE [BLEND {ZMIN | ZMAX}] rmin \ rmax \ tmin \ tmax \ zmin \ zmax \setminus M \ P \ T \ \alpha \ \beta \ [val\_k \ [val\_om]] [sp_1 \ sp_2 \ \ldots sp_n]
```

This keyword is analogous to the XYZ_RANGE keyword, but allows the bounding box to be specified in cylindrical coordinates. It's assumed that the cylindrical z axis is the same as the Cartesian x axis. The bounding box coordinates are:

```
rmin, rmax r_{min} and r_{max} coordinates of bounding box tmin, tmax \theta_{min} and \theta_{max} coordinates of bounding box zmin, zmax z_{min} and z_{max} coordinates of bounding box
```

The remaining input is the same as for XYZ_RANGE.

If BLEND ZMIN or BLEND ZMAX is specified, the initial conditions will smoothly transition from the specified values at z_{min} to freestream values at z_{max} , or from the specified values at z_{max} to freestream values at z_{min} , respectively. If BLEND is not specified, the initial conditions will be uniform, at the specified values.

```
TURBULENT [INFLOW] [MAGNITUDE] vpmag {FPS|MEAN|PERCENT} [SIZE nsiz] \ [SEED seed] [PEAK_K kpeak [PER_FOOT]] [BL_HEIGHT blhgt]
```

For structured grids, this keyword allows the user to specify an unsteady pseudo-turbulent flow at the inflow boundary. It currently only affects the boundary conditions for the mean flow, not the turbulence model equations. Since the mean velocity field typically carries the majority of the energy, this isn't considered critical.

- wpmag Magnitude of the turbulent intensity (i.e., turbulent velocity fluctuations) specified in either feet per second (FPS), fraction of the mean flow velocity (MEAN), or percent of the mean flow velocity (PERCENT). The intensity will be the same in each coordinate direction.
- nsiz Number of grid points on each side of the "box" of pseudo-turbulent inflow. I.e., the box will contain nsiz × nsiz points for 2-D cases, and nsiz × nsiz × nsiz points for 3-D cases. The default value is 64.
- seed A random number seed, used to ensure repeatability of a given run. The default value is 987.0.
- kpeak The wave number (based on the box size unless PER_FOOT is specified) where the energy spectrum is to peak. The default value is 8.0.
- blhgt The height, in feet, of the incoming boundary layer. The turbulence at the inflow boundary will be scaled in this region to account for the presence of the boundary layer. The default is $L_{box}/10$, where L_{box} is the length of a side of the cube bounding the arbitrary inflow boundary.

When the TURBULENT keyword is used, total conditions must be held constant, and HOLD_TOTALS is automatically applied.

```
{VORTEX | SOLIDBODY | ROTATESOLID} M_n P T \alpha \beta xc yc zc {dw1 | dw1 dw2 dw3}
```

For structured grids, these keywords may be used to specify uniform inflow conditions with free-vortex or solid-body rotation superimposed. They are only valid for a perfect gas, and cannot be used with CHEMISTRY. Solid-body rotation may be specified on any arbitrary inflow boundary. For free-vortex rotation, however, the arbitrary inflow boundary must be a constant x, y, or z plane, and the center of rotation must lie on that plane.

M_n	Normal component of Mach number
P	Pressure, psi
T	Temperature, °R
lpha,eta	Average angles of attack and yaw, relative to the Cartesian \boldsymbol{x} direction, in degrees
xc, yc, zc	Center of rotation in physical coordinates
For VORTEX,	
dw1	Vortex strength. (See the ACTUATOR keyword.)

For SOLIDBODY and ROTATESOLID,

The rotational velocity components are added in such a way that total pressure and total temperature are held constant at the inflow boundary. Thus, the TOTAL option should always be used for this mode, since with the STATIC option the computed static pressure and temperature at the inflow boundary may differ from the specified values.

For calculations in a rotating reference frame (see the ROTATE keyword):

- If the SOLIDBODY option is used, the total conditions specified are those in the rotating frame, and are held fixed in that frame. The total conditions in the inertial frame will vary.
- If the ROTATESOLID option is used, the total conditions specified are those in the inertial frame, and are held fixed in that frame. The total conditions in the rotating frame will vary. Note that the ROTATESOLID option is only valid for calculations in a rotating reference frame.

For both free-vortex and solid-body rotation, the flowfield must already be initialized. (See Section 3.6.) I.e., there must be a pre-existing .cfl file. The VORTEX, SOLIDBODY, and ROTATESOLID options cannot be used during a "cold" start.

Special Capabilities

The following arbitrary inflow keywords are only valid for structured grids, and are only applied at the i = 1 computational plane.

```
USERSPEC fs bl1 bl2 npts
y1 \ M \ P \ T \ \alpha \ \beta
y2 \ M \ P \ T \ \alpha \ \beta
\dots
yn \ M \ P \ T \ \alpha \ \beta
```

This option allows the user to specify a 1-D profile normal to the surface, translated through some buttline range, below the vehicle. These conditions will be set last and thus the data will overwrite UNIFORM conditions over the range of interest.

```
Fuselage station of the profile (to be checked against the grid i=1 fuselage station) bl1,\ bl2 Minimum and maximum buttline over which to translate the profile npts Number of points defining the profile y1-yn Normal distance from the wall M Mach number P Pressure, psi P Temperature, P Rappers of attack and yaw, relative to the Cartesian P direction, in degrees
```

The pressure and temperature may be the total or static conditions, depending upon the current setting of the TOTAL/STATIC keyword. If neither STATIC nor TOTAL have been specified under ARBITRARY INFLOW, then the existing switch from the global input parameters is used (default: TOTAL).

One profile can be specified for each zone. There can be 100 points in each profile. The normal distance is always assumed to be from j = 1 (the reference wall is assumed to be at j = 1). bl1 is the minimum buttline and bl2 is the maximum buttline.

```
See Also: TEST 157
```

By default, USERSPEC only specifies conditions below a vehicle. That is, the wall (j = 1) must be above (higher y) the interior grid points. TEST 157 specifies that all points within the specified buttline range will be affected, above and below the vehicle. This should be the default, but isn't.

```
USERCHEM fs bl1 bl2 npts
y1 \ M \ P \ T \ \alpha \ \beta
sp_1 \ sp_2 \dots sp_n
y2 \ M \ P \ T \ \alpha \ \beta
sp_1 \ sp_2 \dots sp_n
\dots
yn \ M \ P \ T \ \alpha \ \beta
sp_1 \ sp_2 \dots sp_n
```

The USERCHEM option is identical to the USERSPEC option, except that chemistry species mass fractions sp_1, sp_2, \ldots, sp_n are added. The order for the mass fractions must be consistent with the order in which the species are listed with the SPECIES keyword in the CHEMISTRY keyword block. Test options can then be set to model the mixing of gas streams which have different chemical compositions. At this time, only mixing can be modeled. The gas streams cannot chemically react.

Note: Only the STATIC input mode is available for chemistry.

Part of an example USERCHEM input block follows. The file sets up a rectangular jet where the jet composition is a mixture of O₂, CO₂, H₂O, NO₂, and N₂.

```
ARBITRARY INFLOW

ZONE 1

USERCHEM 0.0 -10.0 10.0 6
```

```
0.0 0.3 5.70
                    433.1 0.0 0.0
        0.234 0.0
                    0.0
                          0.0 0.766
  123.05 0.3 5.70
                    433.1 0.0 0.0
        0.234 0.0
                    0.0
                          0.0 0.766
  123.05 1.8 5.66 1940.0 0.0 0.0
        0.096
              0.120 0.048 0.0 0.736
  133.75 1.8 5.66 1940.0 0.0 0.0
        0.096 0.120 0.048 0.0 0.736
  133.75 0.3 5.70
                    433.1
                          0.0 0.0
        0.234 0.0
                    0.0
                          0.0 0.766
  257.0 0.3 5.70
                    433.1 0.0 0.0
       0.234 0.0
                    0.0
                          0.0 0.766
ENDINFLOW
TEST 157 1
             USERSPEC ABOVE AND BELOW VEHICLE
```

```
USERKE fs bl1 bl2 npts
y1 \ M \ P \ T \ \alpha \ \beta \ val\_k \ [val\_om]
y2 \ M \ P \ T \ \alpha \ \beta \ val\_k \ [val\_om]
\dots
yn \ M \ P \ T \ \alpha \ \beta \ val\_k \ [val\_om]
```

The USERKE option is identical to the USERSPEC option, except that the values val_k and val_om are added to specify inflow turbulence levels when the SST turbulence model is being used. The various options are described above under the UNIFORM keyword.

```
USERCHEMKE fs bl1 bl2 npts
y1 \ M \ P \ T \ \alpha \ \beta
sp_1 \ sp_2 \ \dots sp_n \ val\_k \ [val\_om]
y2 \ M \ P \ T \ \alpha \ \beta
sp_1 \ sp_2 \ \dots sp_n \ val\_k \ [val\_om]
\dots
yn \ M \ P \ T \ \alpha \ \beta
sp_1 \ sp_2 \ \dots sp_n \ val\_k \ [val\_om]
```

The USERCHEMKE option is identical to the USERCHEM option, except that the values val_k and val_om are added to specify inflow turbulence levels when the SST turbulence model is being used. The various options are described above under the UNIFORM keyword.

See Also: EXTRAPOLATE, INITIALIZE, REINITIALIZE, ROTATE, TURBULENCE

Unstructured Grids

Control Functions

ENDINFLOW

This optional keyword may be used to end the arbitrary inflow definition.

STATIC | TOTAL

Wind-US User's Guide

Arbitrary inflow conditions specified after this keyword will be considered as static or total, depending which of these is set. By default, the flag is set to the value from the FREESTREAM keyword.

With unstructured grids, TOTAL may only be used for an ideal gas.

HOLD_TOTALS

For points with subsonic inflow, specifying HOLD_TOTALS indicates that total temperature, total pressure, and the local flow angles are to be held at their specified values, and the Mach number will be extrapolated. If HOLD_TOTALS is not specified, total temperature, Mach number, and the local flow angles will be held at their specified values, and the total pressure will be extrapolated.

For points with supersonic inflow, all conditions are held at their specified values, whether HOLD_TOTALS is specified or not.

Note that the HOLD_CHARACTERISTICS option is not available for unstructured grids.

For unstructured grids, HOLD_TOTALS is only valid for a perfect gas. The option will be applied at all the arbitrary inflow regions in the zone, and will remain in effect for all following ZONE keywords.

HOLD_TOTALS only works in conjunction with the USURFACE keyword.

```
DIRECTION {SPECIFIED | CYLINDRICAL [X|Y|Z] | NORMAL [TO INFLOW PLANE]}
```

The DIRECTION keyword indicates how the flow angle is to be set at an inflow plane. The options and input are the same as described above for the use of the DIRECTION keyword with structured grids, except that the ALONG [GRID LINES] option does not apply to unstructured grids.

ZONE n

This keyword, which *must* be specified, identifies the zone for which inflow conditions are being set. For example, if zone 2 is an internal jet, conditions other than freestream may be desired at the inflow to zone 2.

Within the ARBITRARY INFLOW keyword block, the ZONE keyword must come *before* any "Condition Specification" keyword for that zone, but *after* any HOLD_TOTALS or DIRECTION keyword for that zone. In addition, inflow conditions may only be specified for one zone at a time.

Condition Specification

UNIFORM
$$[M \ P \ T \ \alpha \ \beta]$$
 $[sp_1 \ sp_2 \ \dots sp_n]$

This keyword is used to specify uniform flow at arbitrary inflow boundaries, at the flow conditions listed below. If the flow conditions are omitted, those specified with the FREESTREAM keyword are used.

M	Mach number
P	Pressure, psi
T	Temperature, ${}^{\circ}R$
α . β	Angles of attack and vaw, relative to the Cartesian x direction, in degrees

 sp_1, sp_2, \ldots, sp_n Species mass fractions. These are required for real gas flows. The order for these values must be consistent with the order in which the species are listed with the SPECIES keyword in the CHEMISTRY keyword block.

Pressure and temperature are static or total, depending on whether STATIC or TOTAL is specified.

```
USURFACE [FROZEN] surface\ M\ P\ T\ \alpha\ \beta\ [sp_1\ sp_2\ \dots sp_n]
```

This keyword is used with unstructured grids to specify uniform flow at arbitrary inflow boundaries, at the flow conditions listed below. The flow conditions are specified in terms of Mach number, pressure, and temperature. Note that unlike the UNIFORM keyword, the flow conditions must be specified.

surface	Surface ID number
M	Mach number
P	Pressure, psi
T	Temperature, ${}^{\circ}R$
α,eta	Angles of attack and yaw, relative to the Cartesian x direction, in degrees
sp_1, sp_2, \ldots, sp_n	Species mass fractions. These are required for real gas flows. The order for these values must be consistent with the order in which the species are listed with the SPECIES keyword in the CHEMISTRY keyword block. Note that unlike the UNIFORM and IJK_RANGE keywords, here the mass fractions are on the same line as the rest of the flow conditions.

The pressure and temperature listed above may be either static or total, depending on whether STATIC or TOTAL is specified.

The FROZEN option may be specified to freeze the inflow conditions on the surface at the specified values. Note that unlike the FROZEN option available with IJK_RANGE for structured grids, here the values specified with the keyword are used, not those in the .cfl file.

The total number of USURFACE surfaces, and IJK_RANGE, XYZ_RANGE, and RTZ_RANGE regions, is limited to 4000.

```
{\tt UNSTEADY}\ \ var\_name\ freq\ ampl\ phase
```

This keyword allows the user to specify unsteady arbitrary inflow conditions. It must be used with (and follow) the USURFACE keyword. Up to ten different perturbations to the inflow conditions may be specified and will be superimposed to create unsteadiness centered about the conditions given with the USURFACE keyword.

var_name	One of the keywords MACH, PRESSURE, TEMPERATURE, ALPHA, BETA, or VELOCITY
freq	Frequency of the perturbation in Hertz
ampl	Amplitude of the perturbation in appropriate variable units
phase	Phase angle of the perturbation in degrees

UNSTEADY may only be used when a constant time step is specified with the CFL# keyword, and only for a perfect gas.

Note that you may specify multiple, independent pairs of USURFACE and UNSTEADY keywords.

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```
XYZ_RANGE [BLEND {XMIN | XMAX}] xmin\ xmax\ ymin\ ymax\ zmin\ zmax \setminus M\ P\ T\ \alpha\ \beta\ [val\_k\ [val\_om]] [sp_1\ sp_2\ \dots sp_n]
```

This keyword is intended for use during the flowfield initialization process (Section 3.6), and allows specification of initial conditions in a specified Cartesian bounding box. The options and input are the same as described above for the use of the XYZ_RANGE keyword with structured grids.

```
RTZ_RANGE [BLEND {ZMIN | ZMAX}] rmin\ rmax\ tmin\ tmax\ zmin\ zmax \ M\ P\ T\ \alpha\ \beta\ [val\_k\ [val\_om]] [sp_1\ sp_2\ \dots sp_n]
```

This keyword is analogous to the XYZ_RANGE keyword, but allows the bounding box to be specified in cylindrical coordinates. The options and input are the same as described above for the use of the RTZ_RANGE keyword with structured grids.

Example

```
ARBITRARY INFLOW
USURFACE 9 1.1 100. 900. 10. 0.
ENDINFLOW
```

AXISYMMETRIC | AXI-SYM — Axisymmetric flow

Structured Grids

{AXISYMMETRIC | AXI-SYM} $yc\ theta$

This keyword allows a two-dimensional structured grid to be run assuming axisymmetric flow. Note that k_{max} must be 1, and that the .cgd file is unaware that the grid is axisymmetric or 2D; that is determined at Wind-US run time.

The user-specified values are defined as:

yc Waterline location of rotation axis, in input units (scaled in Wind-US by XLONG*SCAFAC and offset by YNOSE)

theta Degrees of rotation assumed for metrics. (Changing theta should not affect the solution, but will affect convergence. I like 5 degrees.)

BL_INIT — Boundary layer initialization

Structured Grids

```
\verb|BL_INIT| nb ixs str igrs [delt] nbli nzone
```

For ideal gases (i.e., no CHEMISTRY) and structured grids, the BL_INIT keyword may be used to initialize a j or k boundary with a laminar or turbulent flat plate boundary layer. The user-specified values are defined as:

nb A number from 1 to 4 indicating which boundary to initialize

- 1 for j = 1
- 2 for $j = j_{max}$
- 3 for k = 1
- 4 for $k = k_{max}$

ixs Integer indicating method for specifying the axial location to start the boundary layer:

- 1 to specify the i index
- 2 to specify the x station

str i index or x location to start the boundary layer, depending on the value of ixs

igrs Integer indicating method for specifying the initial height of the boundary layer:

- 1 to specify the number of grid points from the wall
- 2 to specify the thickness
- 3 to compute thickness from initial x station

delt Number of grid points in boundary layer (if igrs = 1), or boundary layer thickness (inches) (if igrs = 2)

nbli Integer indicating type of boundary layer and whether to keep constant thickness or grow with x

- 1 Turbulent profile, growing with x
- -1 Turbulent profile, constant thickness
- 2 Laminar profile, growing with x
- -2 Laminar profile, constant thickness

nzone Zone number for initialization

- 0 All zones
- > 0 Zone nzone only

A flat plate temperature distribution will also be constructed consistent with the wall temperature boundary condition of either adiabatic or constant temperature wall, as specified using the WALL TEMPERATURE keyword. However, this keyword cannot be used with WALL TEMPERATURE EQUILIBRIUM.

Note: BL_INIT cannot currently do more than one boundary in each zone without over-writing itself.

BLEED — Bleed region flow rate

Structured Grids

```
BLEED {MASS region mdot | POROSITY region pplen\ por\ [dis] | \ EMPIRICAL region mode\ blv1\ blv2\ blv3\ blv4 | FORCING region amp\ freq\ phase | \ WALL region | AEDC region dstar\ por\ aspect}
```

Unstructured Grids

```
BLEED {MASS region\ mdot\ |\ POROSITY\ region\ pplen\ por\ dis\ |\ FORCING\ region\ amp\ freq\ phase\ |\ \backslash\ WALL\ region}
```

The effect of bleed on the flow can be modeled, if bleed regions were identified in the grid file. The parameters discussed below identify the bleed rate for each region, for a specific solution. If a bleed region is not named in this file, its bleed rate is set to zero.

There are five possible bleed modes for structured grids, and three for unstructured grids, as described below. Unless noted otherwise, the keywords apply to both structured and unstructured grids.

Bleed is only allowed for perfect gases, for both structured and unstructured grids. In addition, with unstructured grids bleed is only allowed with non-rotating grids. With unstructured grids, the bleed boundary condition is applied at the cell faces, but the flow field values written to the .cfl file for post-processing are at the nodes. The results may thus be slightly different around the edges of the bleed region with structured and unstructured grids.

```
BLEED MASS region \ mdot
```

region Bleed region number from .cgd file

mdot Normalized bleed flow rate

mdot can also be thought of as the mass flow ratio for the bleed region. The actual bleed mass flow is calculated as

$$\dot{m}_b = mdot(\rho_\infty U_\infty A_c)$$

where A_c is an arbitrarily set reference "capture area" that you must specify, in either GMAN (using the CAPTURE AREA command, or the BOUNDARY COND. menu) or MADCAP (using "Set Capture Area" from the "Boundary Conditions" menu), in each zone of the grid file that contains bleed areas.

The bleed velocity will automatically be limited to Mach 1.

Although this is intended as a bleed model, it can also be used for blowing by setting mdot to a negative value.

```
BLEED POROSITY region pplen por dis
```

[There are some questions about the coding for the porous bleed model that need to be resolved. This model should be therefore used with caution.]

region Bleed region number from .cgd file

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pplenBack pressure p_{plen} , in psia

por Porosity

Discharge coefficient; may be defaulted for structured grids dis

With this model, the velocity at the wall will be computed from the local pressure p in the flow field, and the specified back pressure p_{plen} . If $p > p_{plen}$, the flow will be out of the computational domain (i.e., bleed). If $p < p_{plen}$, the flow will be into the computational domain (i.e., blowing).

For unstructured grids, the discharge coefficient must be specified. For structured grids, however, it may be omitted. In this case, for bleed a default value is computed from the specified back pressure and the local flow conditions, using the empirically-based method of Dittrich and Graves (1956). For blowing, the default value for the discharge coefficient is 0.6.

BLEED EMPIRICAL region mode blv1 blv2 blv3 blv4

This keyword specifies use of the empirical bleed model of Mayer and Paynter (1994).

The input parameter region is the bleed region number from the .cqd file. The input data for the bleed model is given by the values of blv1 through blv4. Various combinations of values may be specified, depending on the *mode*, as described below. Modes 1 and 2 are only available for structured grids.

\underline{mode}	blv1	$\underline{blv2}$	blv3	$\underline{blv4}$
1	p_{plen}	Porosity	qsmode	N_{bl}
2	p_{plen}	Porosity	qsmode	
3	p_{plen}	Porosity	qsmode	M
4	Q_{sonic}	Porosity	M	
5	p_{plen}	Porosity		
6	p_{exit}	Porosity	C_{dis}	A_{exit}

In the above table, p_{plen} is the bleed plenum static pressure, N_{bl} is the number of grid points in the boundary layer, Q_{sonic} is the sonic mass flow coefficient (described below), M is the local Mach number at the edge of the boundary layer, p_{exit} is the bleed exit static pressure, p_{exit} is the bleed discharge coefficient, and A_{exit} is the bleed exit area. The parameter qsmode is an integer from 1 to 3 defining how Q_{sonic} is to be computed, as follows:

- Set $Q_{sonic} = 1$
- 2 Compute Q_{sonic} for 90° holes
- Compute Q_{sonic} for 20° holes

In the Mayer-Paynter model, the bleed velocity is given by the formula

$$V_{bleed} = Q_{sonic} \, \Phi \, \Gamma \, T \, \frac{p_T}{p} \, \frac{1}{\sqrt{T_T}}$$

where Φ is the porosity, p and T are the local static pressure and temperature, and p_T and T_T are the local total pressure and temperature. "Local" means at the edge of the boundary layer at the location of the grid point within the bleed region. The parameter Γ is a function of the specific heat ratio γ .

$$\Gamma = \left(1 + \frac{\gamma - 1}{2}\right)^{-\frac{\gamma + 1}{2(\gamma - 1)}}$$

Central to the model is Q_{sonic} , the sonic mass flow coefficient, defined as

$$Q_{sonic} = \frac{\dot{m}_{bleed}}{\dot{m}_{max}} = f\left(\alpha, M, \frac{p_{plen}}{p_T}\right)$$

where \dot{m}_{bleed} is the actual bleed flow rate and \dot{m}_{max} is the maximum theoretical bleed flow rate.

 Q_{sonic} is a function of the bleed hole angle α , the local Mach number M, and the ratio of the plenum pressure p_{plen} to the local pressure p. The functional relationship is in the form of tabulated experimental data for circular bleed holes at angles of 20° (McLafferty and Ranard, 1958) and 90° (Syberg and Hickox, 1972).

BLEED FORCING region amp freq phase

This mode allows an oscillating normal velocity bleed boundary condition to be specified.

region Bleed region number from .cgd file

amp Amplitude of the normal velocity oscillation (ft/sec)

freq Frequency of the oscillation (Hz)

phase Phase offset of the oscillation (deg)

BLEED FORCING was designed to be used on its own, independently of the other bleed modes. However, when BLEED FORCING is used in the same bleed region as another BLEED keyword, the specified oscillatory bleed velocity will be added to the bleed velocity computed for the other bleed mode.

${\tt BLEED} \ {\tt WALL} \ region$

region Bleed region number from .cqd file

This keyword may be used to explicitly turn bleed off in a specific bleed region, and to treat the boundary as a viscous solid wall.

BLEED AEDC region dstar por aspect

region Bleed region number from .cgd file

dstar Displacement thickness at start of bleed region (in)

por Porosity

aspect Bleed hole aspect ratio (i.e., ratio of bleed hole diameter to wall thickness)

This bleed mode is only available for structured grids.

When the AEDC keyword is specified, the bleed region uses the AEDC wind tunnel wall correction correlations. This model is based on the work of Martin, Sickles, and Stanley (Martin, Sickles, and Stanley, 1993), is hardwired for AEDC wind tunnels, and is not intended for general use. It assumes the following:

- Tunnel flow is in the +i direction.
- The bleed region is on a j or k boundary.
- The bleed region encompasses the entire zonal boundary.

See Also: BLOW, MASS FLOW, TEST 46, TEST 67

BLOW — Inject vectored flow over a selected region

Porous wall cooling over a selected region can be simulated using this keyword. The region must be identified as a bleed region in the grid file. This option is intended for mass inflow only (i.e., $P_t > \text{local } P_s$). It won't work well for grids that are skewed at the wall, resulting in blowing mass flow errors.

Blowing may also be modeled using the first two forms of the BLEED keyword (i.e., BLEED and BLEED POROSITY).

The definition of the blowing direction is general enough that a blowing region on the upper and/or lower surface of a wing will be treated consistently with a single specification. However, one should avoid specifying a blowing region in more severe cases, such as the normal part of a backward-facing step, where the surface normal is purely in the x direction.

Except for BLOW MASS, BLOW SURFACE, and BLOW VALVE, blowing is only allowed for perfect gases, for both structured and unstructured grids. In addition, with unstructured grids blowing is only allowed with non-rotating grids. With unstructured grids, the blowing boundary condition is applied at the cell faces, but the flow field values written to the .cfl file for post-processing are at the nodes. The results may thus be slightly different around the edges of the blowing region with structured and unstructured grids.

```
BLOW MASS region\ mdot\ T_{inj} [STATIC|TOTAL] \ angle\_inc\ [ABOUT\_Z\ [angle\_azi]\ |\ ABOUT\_NORMAL\ [angle\_azi]\ ] [BLOW SPECIES_FRACTIONS region\ sp_1\ sp_2\ \dots sp_n] [BLOW FORCING region\ MASS|VELOCITY\ amp\ freq\ phase]
```

region	Bleed region number from $.cgd$ file
mdot	Injected mass flow in region $region$ (lb_m/sec)
T_{inj}	Temperature of injected flow (°R). Either static or total temperature may be specified, as indicated by the optional choice of STATIC or TOTAL. (However, TOTAL may not be used when VELOCITY is used with BLOW FORCING, or for non-perfect gases.) The default is STATIC.
$angle_inc$	Blowing inclination angle (degrees); must be > 0
$angle_azi$	Blowing azimuthal angle (degrees). The default is 0.0.
$sp_1 sp_2 \dots sp_n$	Mass fractions of injected species. The default values are those specified in the CHEMISTRY keyword block.

The blowing direction is set by the input inclination and azimuthal angles.

The inclination angle is the angle between the blowing direction and the projection of the x axis onto the surface.

If ABOUT_Z is specified, the azimuthal angle is the angle between the blowing direction and the projection of the surface normal onto a constant z plane. Starting from the projection of the x axis onto the surface, the blowing direction is thus determined by rotating about the projection of the surface normal onto a constant z plane by the azimuthal angle, then "up" from the surface by the inclination angle.

If ABOUT_NORMAL is specified, the azimuthal angle is the angle between the blowing direction and the surface normal itself. Again starting from the projection of the x axis onto the surface, the blowing direction is thus determined by rotating about the surface normal by the azimuthal angle, then "up" from the surface by the inclination angle.

If neither ABOUT_Z nor ABOUT_NORMAL is specified, ABOUT_Z is assumed, and angle_azi will have its default value of 0.0.

For finite-rate chemistry, when BLOW SPECIES_FRACTIONS is specified the mass fractions should be specified in the same order as the species in the chemistry data (.chm) file, described in Section 7.11. However, if the species have been re-ordered using the SPECIES keyword in the CHEMISTRY keyword block, and the BLOW SPECIES_FRACTIONS keyword comes after the CHEMISTRY keyword block in the input data (.dat) file, then the mass fractions should be specified in the new species order.

BLOW FORCING may be used with this blowing mode to add an oscillatory component to the blowing velocity. The added blowing is specified as either a mass flow or velocity, depending on the choice of MASS or VELOCITY. (If VELOCITY is used, the static temperature must be specified with the BLOW keyword.) The direction of the added blowing will be the same as the mean blowing, as determined by $angle_inc$ and $angle_azi$.

```
 \begin{array}{ll} \textit{region} & \textit{Bleed region number from }.\textit{cgd} \textit{ file} \\ \\ \textit{amp} & \textit{Amplitude of the oscillation for the added blowing (lb_m/sec for MASS, ft/sec for VELOCITY)} \\ \textit{freq} & \textit{Frequency of the oscillation (deg/sec)} \\ \textit{phase} & \textit{Phase offset of the oscillation (deg)} \\ \end{array}
```

```
BLOW PLENUM region\ P_t\ T\ angle [BLOW FORCING region\ MASS\ |\ VELOCITY\ amp\ freq\ phase]
```

```
region Bleed region number from .cgd file P_t Plenum total pressure (psi)

T Plenum static temperature ({}^{\circ}R)

angle Blowing angle relative to x-y plane (degrees)
```

If the flowfield static pressure P_s becomes greater than the plenum total pressure P_t , the plenum total pressure will be automatically increased to $1.005P_s$ to maintain a blowing boundary condition. Setting TEST 52 will notify the user when this occurs.

BLOW FORCING may be used with this blowing mode to add an oscillatory component to the blowing velocity. The added blowing is specified as either a mass flow or velocity, depending on

the choice of MASS or VELOCITY. The direction of the added blowing will be the same as the mean blowing, as determined by *angle*. The input parameters are the same as when BLOW FORCING is used with BLOW MASS, described above.

This blowing mode may only be used with a perfect gas.

```
\begin{tabular}{ll} \it region & \it Region number from .cgd file \\ P_t & \it Plenum total pressure (psi) \\ T_t & \it Plenum total temperature (°R) \\ \it angle\_inc & \it Blowing inclination angle (degrees); must be > 0 \\ \it angle\_azi & \it Blowing azimuthal angle (degrees). The default is 0.0. \\ \end{tabular}
```

If UNSTEADY is specified, unsteady periodic plenum conditions are applied. The plenum conditions over one period must be specified in a table immediately following the BLOW SURFACE keyword, with the following format:

```
PROFILE npts
per(1) amp(1)
per(2) amp(2)
...
per(npts) amp(npts)
```

PROFILE $npts\ time(1)\ press(1)\ temp(1)\ time(2)\ press(2)\ temp(2)\ ...\ time(npts)\ press(npts)\ temp(npts)$ where npts is the number of points in the profile, time is time in seconds, press is total pressure in psi, temp is total temperature in ${}^{\circ}R$.

With BLOW SURFACE, blowing will occur whenever the local flowfield static pressure is less than the specified plenum total pressure. If the flowfield static pressure is greater than the plenum total pressure, the velocity normal to the wall at that point is set to zero (i.e., a solid wall with no blowing or bleed). Unlike the BLOW VALVE capability, with BLOW SURFACE blowing is turned on or off locally, on a point-by-point basis.

The blowing velocity is also constrained to subsonic values.

The blowing direction is set by the input inclination and azimuthal angles. The angles are defined in the same way as when BLOW MASS is used to specify a constant blowing mass flow, as described above.

The BLOW SURFACE keyword may not be used for multi-species flows, but may be used with Liu-Vinokur equilibrium air chemistry.

If the TEST 195 option is set, a message will be written in the list output (.lis) file whenever the blowing is turned off because the flowfield static pressure is too large. Note, however, that this is a five-line message written for each iteration and each "closed" node, and could cause the .lis file to become very large very quickly.

BLOW FORCING may be used with this blowing mode to add an oscillatory component to the blowing velocity. Note that when BLOW FORCING is used with BLOW SURFACE, the added blowing must specified as a mass flow. The VELOCITY option is not available. The direction of the added

blowing will be the same as the mean blowing, as determined by angle_inc and angle_azi. The input parameters are the same as when BLOW FORCING is used with BLOW MASS, described above.²⁹

```
{\tt BLOW~WALL}~region
```

region Bleed region number from .cqd file

This keyword may be used to explicitly turn blowing off in a specific region, and to treat the boundary as a viscous solid wall.

```
BLOW VALVE region\ P_t\ T_t\ angle [BLOW SPECIES_FRACTIONS region\ sp_1\ sp_2\ \dots sp_n]
```

 $\begin{array}{lll} \textit{region} & & \text{Bleed region number from }.\textit{cgd} \text{ file} \\ P_t & & \text{Plenum total pressure (psi)} \\ T_t & & \text{Plenum total temperature (°R)} \\ \textit{angle} & & \text{Blowing angle relative to } \textit{x-y} \text{ plane (degrees)} \\ \textit{sp}_1 \; \textit{sp}_2 \ldots \textit{sp}_n & & \text{Mass fractions of injected species. The default values are those specified in the CHEMISTRY keyword block.} \\ \end{array}$

If the flowfield static pressure P_s becomes greater than the plenum total pressure P_t at any point within the blowing region, blowing will be shut off for the entire region, and the surface will be treated as a solid wall. A *VLV* line is written to the list output (.lis) file whenever the valve changes status.

For finite-rate chemistry, when BLOW SPECIES_FRACTIONS is specified the mass fractions should be specified in the same order as the species in the chemistry data (.chm) file, described in Section 7.11. However, if the species have been re-ordered using the SPECIES keyword in the CHEMISTRY keyword block, and the BLOW SPECIES_FRACTIONS keyword comes after the CHEMISTRY keyword block in the input data (.dat) file, then the mass fractions should be specified in the new species order.

With BLOW VALVE, the blowing region may extend to more than a boundary surface, and may also be split between zones. Note, however, that when a blowing region is split between multiple processors, the separate sub-regions act independently until the end of a cycle. If the flowfield static pressure grows large enough in one sub-region to close the valve but not in the other sub-region(s), the valve will close for the first sub-region, but not on the others until the end of the cycle. The reverse situation (i.e., opening a closed valve) may also occur. This may be prevented by running one iteration per cycle; as a practical matter, it is not expected to cause problems with the default of five iterations per cycle.

See Also: BLEED, TEST 16, TEST 46, TEST 52, TEST 67, TEST 178, TEST 195

²⁹When both BLOW FORCING and UNSTEADY are used with BLOW SURFACE, the forcing is applied after the unsteady plenum conditions. Note, though, that using these together probably doesn't make sense.

${\bf BOUNDARY\text{-}CONDITIONS--Face\ or\ ghost\ cell\ boundary\ conditions}$

Unstructured Grids

BOUNDARY-CONDITIONS {GHOST-CELL | FACE}

The BOUNDARY-CONDITIONS keyword may be used to specify that either face or ghost cell boundary conditions are to be used. The default is face boundary conditions.

This keyword may only be used for perfect gases.

BOUNDARY-DAMP | BDAMP — Boundary damping (block)

Structured Grids

This keyword may be used with structured grids to apply second- and fourth-order smoothing to damp waves in the vicinity of computational boundaries. It is also possible to turn off smoothing near any boundary for any numerical operator (ξ , η , or ζ). This option must be used with the SMOOTHING keyword and TEST 49 2. The user-specified values are defined as:

range Zone specification range, in one of the following forms:

zonenum Selects zone zonenum begzone: endzone Selects all zones from begzone to endzone begzone: Selects all zones from begzone to MAXZONE, the total number of zones in the grid file : endzone Selects all zones from 1 to endzone ALL Selects all zones Selects all zones Starting boundary (0 to begin at boundary 1). The values 1–6 correspond to the i=1, $i=i_{max},\ j=1,\ j=j_{max},\ k=1,$ and $k=k_{max}$ boundaries, respectively. Ending boundary (0 to end at last boundary)

Number of points into the domain over which boundary damping is to be applied

c2 Second-order smoothing coefficient

c4 Fourth-order smoothing coefficient

The NOSMOOTHING keyword is used to turn off smoothing (as specified using the SMOOTHING keyword) for the most recently specified ZONE and BOUNDARY. The keywords NI, NJ, and/or NK specify that smoothing is to be turned off for the ξ , η , and/or ζ operator, respectively, for boundaries bnd1 through bnd2. When NOSMOOTHING is used, at least one of NI, NJ, and NK must be specified.

 m_1, m_2, m_3 Number of points into the domain on boundaries bnd1 through bnd2 over which smoothing is to be turned off

Example

bnd1

bnd2

```
BOUNDARY-DAMP
ZONE 2:4 BOUNDARY 1:3 POINTS 15 SECOND .01 FOURTH .03
ZONE 3:3 BOUNDARY 4:4 POINTS 10 SECOND .02 FOURTH .01
ZONE 5:5
BOUNDARY 2:2
NOSMOOTHING NI 31 NJ 21
END-BOUNDARY-DAMP

See Also: SMOOTHING, TEST 49
```

BOUNDARY TVD — Boundary total variation diminishing operator flag

Structured Grids

```
BOUNDARY TVD [{OFF | FACTOR factor} [ZONE range1[, range2[, . . . rangen]] \ [BOUNDARY {ALL | I1 | IMAX | J1 | JMAX | K1 | KMAX | OVERLAP}]]]
```

This keyword controls the TVD flux limiter in the explicit operator at the specified coupled boundaries in structured grids. The same type of limiter (minmod, Koren, or van Albada) will be used as for the internal scheme.

FACTOR specifies the "compression" parameter for the TVD operator at the specified boundaries, which controls the amount of limiting. Larger numbers result in less limiting towards a first order operator. Setting *factor* to zero is equivalent to a first order operator, one corresponds to no overshoots, and two allows some increase in interface value over surrounding values.

The boundary TVD limiter must be more limiting than the interior TVD limiter for it to have any additional effect. I.e., the value factor must be less than the value used for the internal scheme.

In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

begzone: Selects all zones from begzone to MAXZONE, the total number of zones in the

grid file

: endzone Selects all zones from 1 to endzone

ALL Selects all zones

By default, boundary TVD is on at all boundaries. If the BOUNDARY TVD keyword is not specified at all, the compression parameter will be the same as is used for the internal scheme. If BOUNDARY TVD is specified, but without additional keywords, the default value for *factor* will be used, as listed for the TVD keyword.

Note that if a particular zone (or zones) is to be specified, then either "OFF" or "FACTOR factor" must also be specified. And, if a particular boundary is to be specified, then the zone(s) must also be explicitly specified.

See Also: TVD

CFL# — CFL/time step specification

Structured Grids

```
{CFL# | TIMESTEP} { \
        [MODE 1] [CFL | SECONDS] cfl1 [zone_selector [ityp]] | \
        [MODE 2] [CFL | SECONDS] INCREMENT cfl1 cflmax cflfac inccfl istart \
        [zone_selector [ityp]] | \
        MODE 3 [DT cfl1] [CFLMIN cflmin] [CFLMAX cflmax] [ITIME itime] [zone_selector]}
```

Unstructured Grids

```
{CFL# | TIMESTEP} { \
        [MODE 1] [CFL | SECONDS] cfl1 [zone_selector] | \
        [MODE 2] [CFL | SECONDS] INCREMENT cfl1 cflmax cflfac inccfl istart \
        [zone_selector] | \
        {AUTOMATIC | MODE 4} [CFLSTART cfl] [CFLMIN cflmin] [CFLMAX cflmax] \
        [INCREASE incrfac] [DECREASE decrfac] [FREQUENCY freq] \
        [TRY_INCREASE tryincr] [zone_selector]}
```

This keyword allows the user to specify the CFL number or time step for all zones, or on a zone-by-zone basis. All keywords and parameters must be on one line in the data file. If this keyword is not used, a constant CFL number is used, with a value of 1.3 for structured grids and 10.0 for unstructured grids. For unstructured grids, a larger CFL value can usually be used than for structured grids. Suggested values for unstructured grids are 10 at the start of a calculation, and as high as 50 for later runs.

Structured Grids

For structured grids there are three different options, as specified by the MODE value. Note that the MODE keyword is optional for modes 1 and 2, but required for mode 3. (Modes 1 and 2 are distinguished in the code by whether or not the word INCREMENT is present.)

```
{CFL# | TIMESTEP} [MODE 1] [CFL | SECONDS] cfl1 [zone\_selector [ityp]]
```

In this mode, the CFL number or time step is specified directly, by the input value cfl1.

CFL A CFL number is being specified. This is the default, and should be used for steady flow problems.

SECONDS The time step is being specified in seconds. This option is generally used for unsteady flow problems.

With single-zone grids, when a CFL number is being specified the value of ityp may be set to 1 to indicate that a global time step (i.e., constant in space) should be used, equal to the minimum value in the zone. This allows the time step to be determined through a CFL number for unsteady flow problems. When a CFL number is specified, and ityp = 0 or is omitted, a local time step (i.e., varying in space) will be used.

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{CFL# | TIMESTEP} [MODE 2] [CFL | SECONDS] INCREMENT cfl1 cflmax cflfac incefl istart \ [zone_selector [ityp]]

In this mode, the CFL number or time step will be gradually increased as the calculation proceeds. The CFL and SECONDS options have the same meaning as for mode 1, again with CFL as the default. The increase in the CFL number or time step is controlled by the following input values:

cfl1CFL number or time step for iteration istart

cflmaxMaximum CFL number or time step allowed

cflfacFactor which multiplies the CFL number or time step every *inceft* iterations

Iteration increment at which the CFL number or time step is multiplied by cfffac. The inccflmultiplication is done whenever mod (iteration, inceft) = 1.

Starting iteration in the increment calculation. Note that the iteration values are istartcumulative, starting from the initial run. I.e., to gradually increase the CFL number at the beginning of a restart case, istart should be set to the value of the iteration in the existing .cfl file, from the previous run.

As in mode 1, when a CFL number is being specified for a single-zone grid, the value of ityp may be set to 1 to indicate that a global time step (i.e., constant in space) should be used, equal to the minimum value in the zone.

If mode 2 is used in a staged solution, it should only be used at the start of the run, and SET CFL shouldn't be specified in the SOLVER-STAGES block for the first stage.

{CFL# | TIMESTEP} MODE 3 [DT cfl1] [CFLMIN cflmin] [CFLMAX cflmax] [ITIME itime] \ $[zone_selector]$

This mode uses the time step calculation procedure originally used in the OVERFLOW code.

itimeA flag indicating the type of time step to be used

- Time-accurate run (i.e., time step is constant in space)
- 1 Time step scaled by local metric Jacobian, with fudge factor of 0.005 to keep the time step from getting too small for really tiny cells
- 2 Time step scaled by local metric Jacobian, without a fudge factor
- 3 Constant CFL number equal to cflmax

Time step parameter. For itime = 0 (i.e., time-accurate cases), this is the time step in seconds nondimensionalized by L_r/a_r , where L_r is the grid reference length and a_r is the freestream speed of sound. For itime = 1 and 2, this is a

CFL number, before scaling by the local metric Jacobian. For itime = 3, cfl = 1

is not used. The default value is 0.5.

For itime = 3, the CFL number is set to cflmax, and cflmin is not used. The cflmin, cflmax CFL number is defined using the sum of the maximum eigenvalues in each coordinate direction.

> For other itime values, cflmin and cflmax are used to limit the CFL number to values within the specified range. A value of 0.0 indicates no limiting. If either cflmin or cflmax is negative, the absolute values are used, and the CFL number is defined using the method of Gnoffo, with a viscous correction by Tannehill. If

cfl1

both *cflmin* and *cflmax* are non-negative, the usual one-dimensional (inviscid) CFL number definition using the maximum eigenvalue is used.

The default values are both 0.0.

If mode 3 is used, it must be used in all zones, but the values of cf11, cflmin, etc., may vary from zone to zone.

See Also: CROSSFLOW, NEWTON, TEMPORAL

Unstructured Grids

There are also three options for specifying the CFL number or time step for unstructured grids. Note that the MODE keyword is optional for the first two options, but the word AUTOMATIC (or optionally MODE 4), is required for the third option.

```
{CFL# | TIMESTEP} [MODE 1] [CFL | SECONDS] cfl1 [zone\_selector]
```

This mode is essentially the same as the MODE 1 option for structured grids, described above. However, with unstructured grids the *ityp* input option, which may be used with single-zone structured grids when a CFL number is being specified to indicate that a global time step (i.e., constant in space) should be used, is not available. In addition, for unstructured grids with local time stepping, the size of the physical time step will automatically be limited to the largest time step in the grid that corresponds to a CFL number of one. Of course, due to variations in grid spacing and flow conditions, the CFL number at other locations may be much higher.

```
{CFL# | TIMESTEP} [MODE 2] [CFL | SECONDS] INCREMENT cfl1 cflmax cflfac incefl istart \setminus [zone\_selector]
```

This mode is essentially the same as the MODE 2 option for structured grids, described above. Like MODE 1, with unstructured grids the *ityp* input option is not available, and with local time stepping, the size of the physical time step will automatically be limited to the largest time step in the grid that corresponds to a CFL number of one.

```
{CFL# | TIMESTEP} {AUTOMATIC | MODE 4} [CFLSTART cfl] [CFLMIN cflmin] \ [CFLMAX cflmax] [INCREASE incrfac] [DECREASE decrfac] [FREQUENCY freq] \ [TRY_INCREASE tryincr] [zone\_selector]
```

With this mode, the CFL number is automatically decreased or increased based on the convergence behavior of the solution. Note that like all the other modes, the keyword may be specified as CFL# or TIMESTEP, but this mode specifies only a CFL number, not a physical time step.

The input values are:

```
cfl1 Starting CFL number. The default is 10.0.
```

cflmin Minimum CFL number that will be used. The default is 0.1.

cflmax Maximum CFL number that will be used. The default is $100 \times cfl11$.

incrfac Factor by which the CFL number will be increased in a non-diverging solution. The default is 2.0.

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decrfac Factor by which the CFL number will be decreased in a diverging solution. The default

is 10.0.

freq Iteration frequency for checking whether or not the CFL number should be increased in a non-diverging solution. This is referenced to the last iteration in which the CFL number was changed (increased or decreased). The default is 10.

tryincr Interval for increasing the CFL number in a solution that is neither converging or diverging. The default is 3.

The solution is checked for divergence every iteration, and if it is diverging, the CFL number is decreased by the factor decrfac, down to a minimum of cflmin.

In a non-diverging solution, the solution is checked every freq iterations to determine whether or not the CFL number should be increased. If the solution is converging, the CFL number is increased by the factor incrfac. If the solution is neither converging or diverging, the CFL number is also increased by the factor incrfac, but only every tryincr times this check is made (i.e., every freq × tryincr iterations). The CFL number in a particular zone is limited to the value of cflmax for that zone, or 5 times the smallest CFL number for all the other zones, whichever is lower.

The $\Delta \mathbf{Q}$ limiter (keyword DQ LIMITER) must be on (which is the default) when CFL AUTOMATIC is used.

See Also: NEWTON, TEMPORAL, DQ LIMITER

Examples

Set the CFL number to 1.5 in all zones.

```
CFL# CFL 1.5
```

Set the CFL number to 1.5 in zones 1 and 5, and 0.7 in zones 2, 3, and 4.

```
CFL# CFL 1.5 ZONE 1,5
CFL# CFL 0.7 ZONE 2:4
```

The following example sets the CFL number to 1.0, computes the corresponding time step at every point in the grid, finds the minimum of those time steps, and resets the time step at every point to that minimum value. This allows the time step to be set for an unsteady flow problem by specifying a CFL number. Note that even though this may only be done for single-zone grids, the syntax requires that the zone be explicitly specified.

```
CFL# CFL 1.0 ZONE 1 1
```

Set the time step equal to 1×10^{-6} seconds at every point.

```
CFL# SECONDS 0.000001
```

The next example sets the CFL number to 0.5 for the first 500 iterations, then increases it by a factor of 1.1 every 100 iterations. The maximum value allowed will be 2.0, so the actual final CFL number will be some value just below 2.0. (For this example, it's 1.899.)

```
CFL# CFL INCREMENT 0.5 2.0 1.1 100 501
```

The remaining examples use the mode 3 time step calculation procedure, only available for structured grids. The first example sets the time step parameter to 0.1, and defaults the remaining input parameters, causing the actual time step to be scaled by the local metric Jacobian with the 0.005 fudge factor, with no minimum or maximum value.

```
CFL# MODE 3 DT 0.1
```

In this example the mode 3 time step calculation procedure is used to set the time step parameter to 0.5. The actual time step will be scaled by the local metric Jacobian with the 0.005 fudge factor, with minimum and maximum CFL numbers of 1.0 and 5.0. Since the minimum and maximum values are specified as positive numbers, the standard one-dimensional (inviscid) CFL number definition using the maximum eigenvalue is used.

```
CFL# MODE 3 DT 0.5 CFLMIN 1.0 CFLMAX 5.0
```

The next example uses the mode 3 time step calculation procedure to set the time step parameter to 0.05. The actual time step will be scaled by the local metric Jacobian without the 0.005 fudge factor, with minimum and maximum CFL numbers of 1.0 and 5.0. Since the minimum and maximum values are specified as negative numbers, the CFL definition of Gnoffo is used, with a viscous correction by Tannehill.

```
CFL# MODE 3 DT 0.05 ITIME 2 CFLMIN -1.0 CFLMAX -5.0
```

Use the mode 3 time step calculation procedure to set the CFL number to a constant value of 1.3.

```
CFL# MODE 3 ITIME 3 CFLMAX 1.3
```

Use the mode 3 time step calculation procedure for an unsteady case, with the nondimensional time step set to a constant value of 0.05.

```
CFL# MODE 3 ITIME 0 DT 0.05
```

The following example uses the mode 3 time step calculation procedure for an unsteady case, again with a specified nondimensional time step of 0.05. In this case, however, if the specified time step results in a CFL number at some point in the flow field less than 0.25 or greater than 5.0, the actual time step throughout the flow field will be reset to the corresponding value.

CFL# MODE 3 ITIME 0 DT 0.05 CFLMIN 0.25 CFLMAX 5.0

CGNSBASE — Use CGNS files

CGNSBASE basename

This keyword specifies that the grid and flow data are obtained from, and written to, CGNS (CFD General Notation System) files, not common files. The user-specified basename is the name of the CGNSBase_t node in the grid and flow files.³⁰

It should be noted that, strictly speaking, there is no notion of a CGNS file, only of a CGNS database implemented within one or more ADF (Advanced Data Format) files. We nevertheless use the terminology "CGNS file" in the Wind-US documentation. In addition, a true CGNS database has a single CGNSBase_t node as its root. In the current Wind-US implementation, however, the grid and flow solution are stored in separate files, with both files having their own CGNSBase_t node with the same basename.

When CGNS files are used, the extensions for the grid and flow files must be .cgd and .cfl, respectively, the same as when common files are used. If the CGNSBASE keyword is used, the files are read as CGNS files. If the CGNSBASE keyword is not used, they are assumed to be common files.

Wind-US supports the most common CGNS features, but not all. Some specific limitations on the implementation of CGNS in Wind-US are listed below.

- Many Wind-US input options may be specified on a zonal basis, using the zone number to identify the zone. In the CGNS standard, zones are identified by a name. When CGNS files are used in Wind-US, it is assumed that the zone number maps to the zone's position in an alphanumerically sorted list of zone names.
 - Note that the naming convention Zonen, where n is the zone number, is alphanumeric only up to Zone9. Zone10 through Zone19 would get sorted between Zone1 and Zone2, and so on. However, spaces are allowed in names, so "Zone n", with two spaces, (e.g., Zone 1, Zone 2, ..., Zone 99, Zone100, ...) is alphanumeric up to Zone999.
- The CGNS default names "GridCoordinates" and "FlowSolution" are used for the CGNS GridCoordinates_t and FlowSolution_t nodes. For each zone, only one grid and one flow solution are stored.
- In a CGNS database, the scaling data must be stored with the data itself, but the units to be used may be stored with the data or higher in the node tree. The Wind-US implementation assumes that the units are stored with the data.
- If grid velocities are stored in the grid file, it is assumed that the units are ft/sec.
- The CGNS nodes for storing time-dependent data are not yet supported in Wind-US.
- For overlapping grids, CGNS allows hole points to be identified using multiple OversetHoles_t nodes. Wind-US currently assumes that all hole points in a zone are identified in a single OversetHoles_t node, using a PointList.
- Wind-US writes convergence and time history information to the list output (.lis) and time history (.cth) files, respectively, not to a CGNS file.

There are also some Wind-US features and capabilities that are not supported by the current CGNS standard. These include chemistry and MFD flows, specified boundary layer transition, and bleed boundary conditions. Several proposals are pending to extend the CGNS standard, but for

³⁰Detailed information on the CGNS standard may be found at the CGNS web site, at http://cgns.sourceforge.net/.

now CGNS files created by Wind-US for these types of flows may not be fully CGNS compliant, and should be considered unique to the Wind-US implementation.

CHEMISTRY — Chemistry model selection (block)

```
CHEMISTRY
   {EQUILIBRIUM
       LIU-VINOKUR |
    FROZEN
       [ADJUST [CURVES FOR O K REFERENCE]]
       FILE filename
       SPECIES {AIR | S_1 \ sf_1 \ S_2 \ sf_2 \ \dots S_m \ sf_m}
       [DIFFUSION {NONE | SINGLE | EFFECTIVE-BINARY}]
       [THERMO9] |
   FINITE [RATE]
       [ADJUST [CURVES FOR O K REFERENCE]]
       FILE filename
       SPECIES {AIR | S_1 \ sf_1 \ S_2 \ sf_2 \ \dots S_m \ sf_m}
       [DIFFUSION {NONE | SINGLE | EFFECTIVE-BINARY}]
       [FIXED [ZONE] range1[, range2[, ... rangen]]]
       [FUEL [AIR] [RATIO] value]
       [OMIT [THIRD-BODY] S_1 S_2 ...S_m ZONE range1[, range2[, ... rangen]]]
       [EDC value]
       [DAMKOHLER Da max]}
   [VISCOSITY {SUTHERLAND | WILKE | KEYE | CONSTANT vis | TUNNEL9 | \
                CUSTOM c_1 c_2}]
ENDCHEMISTRY
```

This option allows the user to specify the real gas chemistry mode and input data for the desired species and reactions.

Note: The ARBITRARY INFLOW keyword block, if used, must come after the CHEMISTRY keyword block in the input data (.dat) file.

The various elements of the CHEMISTRY input block are defined as follows:

CHEMISTRY

Defines the beginning of the CHEMISTRY block.

```
{EQUILIBRIUM | FROZEN | FINITE [RATE]}
```

The main part of the CHEMISTRY block is divided into three sections, corresponding to input for equilibrium chemistry, frozen chemistry, and finite-rate chemistry. One and only one of the keywords EQUILIBRIUM, FROZEN, and FINITE RATE must be specified.

```
LIU-VINOKUR
```

Specifies that the Liu and Vinokur equilibrium air curve fits, valid to 50000 K, should be used. Data for the individual species is not output.

This keyword only applies to EQUILIBRIUM chemistry, and must be specified.

ADJUST [CURVES FOR O K REFERENCE]

In the chemistry data (.chm) file, the WINDNASA and NASA3287 curve fits for the thermodynamic properties use an enthalpy reference state of 298.15 K. When the ADJUST keyword is used, the heat of formation at 0 K is read from the .chm file, and the values of enthalpy and Gibb's free energy returned by the WINDNASA and NASA3287 curve fits are shifted to change the reference state to 0 K. If ADJUST is not used, the enthalpy reference state is left as 298.15 K, and the line in the .chm file containing the heat of formation at 0 K, if present, is ignored.

Note that the ADJUST keyword has no effect when the SPARKCRV curve fits are used. They always use an enthalpy reference state of 0 K.

The ADJUST keyword applies to both FROZEN and FINITE RATE chemistry. If specified, it must come before the FILE keyword in the .dat file.

FILE filename

Specifies the chemistry data (.chm) file containing the data defining the thermodynamic property, reaction rates, and transport properties. (See Section 7.11 for the file format.) Unless the chemistry data file is located in the same directory as the one in which Wind-US is running (see the -runroot and -runinplace wind script options) the specified filename must be a full path name. And in either case, it should include the .chm extension.

The FILE keyword applies to both FROZEN and FINITE RATE chemistry, and must be specified.

SPECIES {AIR | S_1 sf_1 S_2 sf_2 ... S_m sf_m }

Specifies the species names $(S_1 \text{ through } S_m)$, and their freestream (reference) mass fractions $(sf_1 \text{ through } sf_m)$. The specified species must be the same as those in the chemistry data (.chm) file, but not necessarily in the same order.

AIR may be specified instead of individual species to indicate use of air mass fractions. The species 02 and N2 must be listed in the chemistry data (.chm) file.

The SPECIES keyword applies to both FROZEN and FINITE RATE chemistry, and must be specified.

DIFFUSION {NONE | SINGLE | EFFECTIVE-BINARY}

Specifies the type of species diffusion. The default is SINGLE, which uses a single mixture diffusion coefficient based on the mixture viscosity and the Schmidt number.

If EFFECTIVE-BINARY is specified, an effective binary diffusion model is used, with each species having its own effective diffusion coefficient with respect to the mixture. (For details see Lankford and Mani (2003).) This option requires that the Lennard-Jones parameters used in computing the intermolecular potential be specified in the chemistry data (.chm) file.

The DIFFUSION keyword applies to both FROZEN and FINITE RATE chemistry.

THERMO9

Simulate AEDC's Tunnel 9 facility, using a real gas table look-up of thermodynamic properties. Either nitrogen or hydrogen may be modeled, as specified by the first species listed in the chemistry

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data (.chm) file. Section 7.11. If the first species name is anything other than N2 or H2, the job will abort.

If THERMO9 is specified, it must come after the FILE keyword in the input data (.dat) file. In addition, if the WINDNASA or NASA3287 format is used for the thermodynamic properties in the .chm file, ADJUST must be specified to shift the enthalpy reference state from 298.15 K to 0 K.

The THERMO9 keyword only applies to FROZEN chemistry, and is only available for structured grids.

```
FIXED [ZONE] range1[,range2[, ... rangen]]
```

Specifies fixed chemistry composition (i.e., a fixed set of mole fractions) in the specified zone(s). Species concentrations are set to the values specified with the SPECIES keyword. This allows the thermodynamic properties to be computed using the correct multi-species models, but the expense of actually solving the species equations is skipped. The chemistry source term is also not computed, since this approach only makes sense where no reactions are taking place.

In the zone specification, the range parameter(s) may be

zonenumSelects zone zonenum

Selects all zones from begzone to endzone begzone: endzone

ALL Selects all zones

This option is intended for use in multi-zone computations with reactions occurring in some zones, but not others. An example is an internal/external nozzle flow, where no reactions take place in the external flow upstream of the nozzle exit. Using this option may significantly lower the execution time, but must be used with care.

The FIXED keyword only applies to FINITE RATE chemistry, and is only available for structured grids.

```
FUEL [AIR] [RATIO] value
```

value specifies the fuel-air ratio for H₂-air combustion (two-reaction global model).

The FUEL AIR RATIO keyword only applies to FINITE RATE chemistry, with ispec = 2 in the chemistry data (.chm) file.

```
OMIT [THIRD-BODY] S_1 S_2 ...S_m ZONE range1[, range2[, ...
```

This keyword allows control of the reacting species in a finite-rate chemistry solution on a zonal basis. Reactions leading to the production or destruction of the species given by S_1 through S_m will be eliminated in the selected zones. In addition, if the THIRD-BODY keyword parameter is used, any reaction involving the named species will be eliminated. The species will still be convected and diffused, maintaining conservation at zone boundaries.

Eliminating reactions in zones where they aren't relevant improves the overall efficiency of the calculation in two ways. First, the time required to perform the unnecessary computations is eliminated. And second, the stiffness of the system of equations is reduced, leading to faster convergence and greater stability.

In the zone specification, the range parameter(s) may be

zonenum Selects zone zonenum begzone: endzone
Selects all zones from begzone to endzone
Selects all zones

Multiple OMIT keywords are allowed, with conflicting input resolved by using the latter of the conflicting entries.

Currently, you cannot omit the final species specified with the SPECIES keyword.

 $\mathtt{EDC}\ value$

Specifies use of the "eddy dissipation concept" when computing the reaction rate for finite-rate chemistry, and sets the value of the eddy dissipation coefficient C_{edc} . See the description of "Format 2" in Section 7.11.3, for more detail about the eddy dissipation concept.

The EDC keyword only applies to FINITE RATE chemistry solutions using the SST turbulence model. It may only be used when ispec = 137 in the chemistry data (.chm) file.

DAMKOHLER Da max

Specifies use of the Damkohler rate limiter when computing the reaction rate for finite-rate chemistry. This is intended to improve the solution convergence rate by reducing the stiffness of the governing equations. Both forward and backward reaction rates will be limited such that the ratio

$$Da = \frac{\tau_{FD}}{\tau_{CH}} \le Da_{max}$$

where τ_{FD} and τ_{CH} are the fluid dynamic and chemical reaction time scales, respectively. A reasonable value for Da_{max} is 10. See the description of "Format 2" in Section 7.11.3, for more detail about the Damkohler rate limiter.

The DAMKOHLER may only be used when ispec = 137 in the chemistry data (.chm) file.

MASS [FRACTIONS] {AIR | $sf_1 \ sf_2 \ \dots sf_n$ }

Specifies the freestream (reference) mass fractions of the species. The mass fractions are specified in the same order as the species in the chemistry data (.chm) file.

AIR may be specified instead of individual species to indicate use of air mass fractions. The species 02 and N2 must be listed in the chemistry data (.chm) file.

The MASS FRACTIONS keyword may be used instead of the SPECIES keyword to specify the species mass fractions, but its use is discouraged. It has been retained for compatibility with input data (.dat) files used with earlier versions of Wind-US.

VISCOSITY {SUTHERLAND | WILKE | KEYE | CONSTANT vis | TUNNEL9 | CUSTOM c_1 c_2 }

Selects the transport property equations. For unstructured grids, only the SUTHERLAND and WILKE options are available. See the VISCOSITY keyword for details.

ENDCHEMISTRY

Defines the end of the CHEMISTRY block.

See Also: GAS, PRANDTL, SCHMIDT, VISCOSITY, DEBUG 16, TEST 5, TEST 66, TEST 69, TEST 70, TEST 71, TEST 88, TEST 89, TEST 90, TEST 91, TEST 92, TEST 93, TEST 94, TEST 95, TEST 96, TEST 99, TEST 164, TEST 172, TEST 178

General and Standard Chemistry Packages

The chemistry data files supplied with Wind-US are summarized in Table 12.31

Table 12: General and Standard Chemistry Packages

Species	File(s)	T_{max}	Nc	Nr	3d
O_2 , NO , O , N , N_2	$air ext{-}5sp ext{-}std ext{-}06k.chm$	6000 K	1	5	V
	$air ext{-} 5sp ext{-} gen ext{-} 06k.chm$				
O_2 , NO , O , N , N_2	$air ext{-}5sp ext{-}std ext{-}15k.chm$	15000 K	3	5	V
	$air ext{-} 5sp ext{-} gen ext{-} 15k.chm$				
O_2 , NO , O , N , N_2	$air ext{-}5sp ext{-}std ext{-}30k.chm$	30000 K	5	5	V
	$air ext{-} 5sp ext{-} gen ext{-} 30k.chm$				
O_2 , NO, O, N, NO ⁺ , e ⁻ , N_2	air-7 sp - std -0 $6k$. chm	6000 K	1	6	V
O_2 , NO, O, N, NO ⁺ , e ⁻ , N_2	air-7 sp - std -15 k . chm	15000 K	3	6	V
	air-7 sp - gen -15 k . chm				
O ₂ , NO, O, N, NO ⁺ , e ⁻ , N ₂	air-7sp-std-30k.chm	30000 K	5	6	V
	air-7 sp - gen -30 k . chm				
O_2 , OH , H_2 , H_2O , N_2	$h2air ext{-}5sp ext{-}std ext{-}06k.chm$	6000 K	1	2	
	$h2air ext{-}5sp ext{-}std ext{-}15k.chm$	15000 K	3		
O_2, H, H_2, H_2O, N_2	Not yet available				
O ₂ , H, H ₂ , H ₂ O, OH, O, N ₂	h2air-7sp-std-15k.chm	15000 K	3	8	
	h2air-7sp-gen-15k.chm				
	h2air- $7sp$ - bak - $15k$. chm				
O_2 , CO_2 , H_2O , NO_2 , N_2	Not yet available				

 $^{3^{1}}$ In Table 12, the symbol e⁻ is electron density, and NO⁺ is ionized nitrogen oxide. Also, T_{max} is the maximum temperature at which the curve fits are valid, Nc is the number of thermodynamic curve fit segments, Nr is the number of reaction equations, and 3d denotes Variable or Average 3rd-body efficiency.

/ — Comment lines

In the extended portion of the file, a "comment line" may be placed anywhere. This line is denoted by a / in column 1. The remainder of the line is disregarded by the input routines, as if the comment line did not exist. The purpose of this option is to allow the user to place explanatory notes within the flow condition data file. A blank line is also ignored.

COMPRESSOR FACE — Outflow boundaries, compressor face

Structured Grids

This keyword allows an outflow boundary in a structured grid to be modeled as a compressor face. In the zone specification, the *range* parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

There are several models available, as described below. All are based on the observation that turbine engine conditions set the corrected mass flow, and that this corresponds directly to the average Mach number at the compressor face. These boundary conditions have been implemented mainly for the analysis of unsteady flow; however, they have also been shown to be robust for the establishment of steady-state, supercritical inlet flows.

```
COMPRESSOR [FACE] [CHUNG] mach [ZONE] range1[,range2[, ... rangen]]
```

The CHUNG keyword option uses the Chung-Cole model (Chung and Cole, 1996). Based on the specified mass-averaged Mach number mach, and using the computed local Mach number and total pressure, a new local static pressure is computed.

Note: This boundary condition is identical in its effect on the flow to the DOWNSTREAM MACH keyword.

The BOEING keyword option uses uses a model developed by Mayer and Paynter (1994). The parameters to be specified as *val1* and *val2* depend on *mode*, as shown in the following table.

\underline{mode}	$\underline{val1}$	$\underline{val2}$
1	Average Mach number	_
2	$ {\rm Corrected\ mass\ flux\ per\ unit\ area,} \\ {\rm lb_m/sec/ft^2} $	_
3	Nominal Mach number	Nominal total temperature, °R

Modes 1 and 2 are equivalent, and are useful in obtaining initial flow fields, and for simulating instantaneous changes at the compressor face via a restart. Mode 3 is used to set a constant volumetric flow condition. Typically, an initial condition is obtained using mode 1 and becomes the nominal condition. Then val1 is set to the Mach number used in mode 1 and val2 is set to the freestream total temperature, i.e., it is assumed that the total temperature at the compressor face is equal to the freestream value. If the average total temperature at the compressor face does not change, then modes 1 and 3 will yield identical results. However, if the average total temperature at the compressor face changes due to a freestream disturbance, for example, then the corrected flow rate will be adjusted to maintain a constant volumetric flow rate.

```
COMPRESSOR [FACE] PAYNTER mach\ val1\ val2 [ZONE] range1[, range2[, ... rangen]
```

The Paynter compressor face model (Paynter, 1998) computes unsteady acoustic reflections from a compressor face subjected to acoustic and convective disturbances. This model is intended to be used for time-accurate simulations of unsteady inlet flow. The CFL number should be less than one. The flow is assumed to be subsonic and axial as it enters the compressor face.

The model assumes the compressor face is positioned within an annular duct with a hub and case, and consists of a single row of axial flow compressor blades. The conditions at the compressor face are determined by the convective velocity response coefficient α , and the acoustic response coefficient β . The definitions of α and β depend on the passage Mach number $M_{passage}$, which is given by

$$M_{passage} = M_{cf}/\cos\Gamma$$

where M_{cf} is the compressor face Mach number, and Γ is the local stagger angle.

For subsonic flow $(M_{passage} < 1)$,

$$\alpha = \frac{\gamma M_{cf}^2}{1-M_{cf}}\tan\left(\frac{\Gamma}{2}\right)\tan\Gamma$$

$$\beta = \tan^2\left(\frac{\Gamma}{2}\right) \left(\frac{1 + M_{cf}}{1 - M_{cf}}\right)$$

where γ is the specific heat ratio. For supersonic flow $(M_{passage} > 1)$,

$$\alpha = \gamma M_{cf}$$

$$\beta = 1$$

Details on the implementation of this boundary condition are given by Slater and Paynter (2000).

In Wind-US, α and β may be treated as constant and specified directly, or computed using the above equations. The local stagger angle Γ is determined by assuming a linear variation between specified values of the stagger angles at the hub and case. The solidity of the blade row is assumed to be greater than one.

The absolute value of the keyword parameter mach sets the nominal or average compressor face Mach number M_{cf} . The meanings of val1 and val2 depend on the sign of mach:

\underline{mach}	$\underline{val1}$	$\underline{val2}$
< 0	Stagger angle (degrees) of the	Stagger angle (degrees) of the
	compressor fan blade at the hub	compressor fan blade at the case
> 0	Constant convective velocity re-	Constant convective velocity re-
	sponse coefficient, α	sponse coefficient, β

```
COMPRESSOR [FACE] SAJBEN mach\ angh\ angc [ZONE] range1[, range2[, ... rangen]]
```

The Sajben compressor face model is basically the same as the Paynter model, except that Sajben's expression for the acoustic response coefficient is used for subsonic flow (Sajben, 1999).

Sajben's expression for the acoustic response coefficient may be written as

$$\beta = \frac{M_{cf} \tan^2 \Gamma}{2(1 + M_{cf}) + M_{cf} \tan^2 \Gamma}$$

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where M_{cf} is the compressor face Mach number, and Γ is the local stagger angle.

With the COMPRESSOR [FACE] SAJBEN keyword, the keyword parameter mach is the nominal or average compressor face Mach number M_{cf} , and angh and angc are the stagger angles in degrees of the compressor fan blade at the hub amd case.

Examples

```
COMPRESSOR FACE CHUNG 0.3 1
COMPRESSOR FACE BOEING 1 0.3 1
COMPRESSOR FACE BOEING 2 0.5516 1
COMPRESSOR FACE BOEING 3 0.3 520.0 1
```

 $See\ Also:$ DOWNSTREAM MACH, DOWNSTREAM PRESSURE, MASS FLOW, TEST 123, TEST 153

CONVERGE — Controls convergence

Structured Grids

```
CONVERGE {LEVEL | ORDER} val
```

Unstructured Grids

```
CONVERGE LOAD [FREQUENCY ncyc] val
```

This keyword allows the user to specify the criterion used to automatically stop a run when it converges.

Note that this keyword should not be used as the sole method for determining convergence. Convergence should be monitored throughout a run by tracking residuals, integrated forces, moments, and mass flow, and/or flow variables at particular locations. See Section 6 for more detail.

Structured Grids

The options available for structured grids are:

LEVEL Check for maximum residual less than val.

ORDER Check for reducing maximum residual by val orders of magnitude.

When the specified convergence level has been reached for all zones, the calculation will stop. The default, when the CONVERGE keyword isn't used, is a four-order of magnitude decrease in the maximum residual.

If Newton iteration is being used, this keyword specifies the convergence criteria within a Newton time step. The overall global convergence criteria is specified using the NEWTON keyword.

See Also: LOADS, HISTORY, NEWTON, TEST 128

Unstructured Grids

For unstructured grids, specifying CONVERGE LOAD tells Wind-US to stop the calculation when the forces and/or moments on user-specified surfaces (and/or subsets of surfaces) have leveled off within a specified tolerance. The LOADS keyword block must also be used to specify the surfaces and/or subsets, and whether forces and/or moments should be computed. The forces and moments used with CONVERGE LOAD will be the sum of the pressure and viscous values, if VISCOUS is included in the surface/subset specification in the LOADS block.

As part of the process, the flux dissipation parameter specified using the EXPLICIT option with the LAX DAMPING keyword, and/or the slope limiting parameter specified using the TVD keyword, are driven to limiting values in each zone as the solution proceeds. With standard and "super-bee" slope limiting, for the Rusanov (Lax) scheme both the flux dissipation and slope limiting parameters are used, while for the Roe scheme, only the slope limiting parameter is used. With Venkatakrishnan slope limiting, for the Rusanov (Lax) scheme only the flux dissipation parameter is used. CONVERGE LOAD can't be used with Venkatakrishnan slope limiting for the Roe scheme.

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For the Rusanov (Lax) scheme, the flux dissipation parameter and the slope limiting parameter (where applicable) are driven to 0.11 (or lower) and 3.49 (or higher), respectively. For the Roe scheme, the slope limiting parameter is driven to 2.49 (or higher).

Convergence is checked every ncyc cycles, with a default of 50. The flux dissipation and/or slope limiting parameters are adjusted when the relative changes in the forces and moments on the specified surfaces within the current zone are all below the value val.

Convergence is assumed when the flux dissipation and/or slope limiting parameters have reached their limits in all zones, and the maximum relative change in the forces and moments, summed over all the zones, is less than 0.01. (The convergence criterion of 0.01 may be changed using DEBUG 24).

See Also: LOADS, LAX DAMPING, TVD, DEBUG 24

COUPLING — Zone coupling mode specification

Structured Grids

COUPLING [MODE] {CHARACTERISTIC | ROE [$\underline{\text{HIGH}}$ | LOW]}

This keyword controls the zone coupling algorithm used for multi-zone solutions. The two possible options are defined below.

CHARACTERISTIC

This coupling algorithm uses one-dimensional characteristic flow theory to set boundary flowfield variables based on local flow direction and strength. These boundary variables are then transferred between zones using interpolation factors stored in the grid file (the .cgd file).

ROE

The Roe coupling algorithm is more consistent with Roe's flux-difference splitting scheme, which is the default explicit operator. Instead of transferring flowfield variables between zones, this algorithm transfers flux cell "interface states," which are critical quantities in Roe's scheme. In this coupling mode, the zone boundary may be thought of as a "cell interface" in Roe's scheme.

The HIGH option, which is the default, turns on higher-order coupling at zone boundaries. With higher-order coupling the solution derivatives are also passed between coupled zones. This results in a slight increase in memory requirements. The derivative information is used to increase the accuracy of the coupling scheme and to improve robustness at coupled boundaries.

The type of coupling that is allowed depends on the type of explicit differencing operator being used, set using the RHS keyword.

- If RHS CENTRAL or RHS COAKLEY is specified, characteristic zone coupling is automatically used.
- Unless COUPLING CHARACTERISTIC is specified, low-order Roe coupling is automatically used for first-order explicit operators.
- Roe coupling (low- or high-order) requires that the Roe, Van Leer, HLLE, HLLC, or Rusanov explicit operator be used.
- High-order Roe coupling may not be used with third-order fully upwind, or fourth- or fifth-order explicit operators.

See Also: RHS

CROSSFLOW — Crossflow CFL factor

Structured Grids

```
{\tt CROSSFLOW} \ \ [{\tt CFL}] \ \ \ [{\tt FACTOR}] \ \ val \ \ [{\tt zone\_selector}]
```

The CROSSFLOW keyword sets the cross-flow CFL factor to val in the specified zones.

For most Wind-US calculations, the time step is based on a constant CFL number throughout the flow field. Thus, for non-uniform grids, the solution advances at different rates in different parts of the grid. Because the CFL number is directly proportional to the physical time step and inversely proportional to the local grid spacing, the physical time step is small where the grid spacing is small, in order to maintain a constant CFL number. For example, in boundary layers and shear layers, where the grid is often closely packed for better resolution, the solution advances more slowly than in other parts of the flowfield. In fact, most of the iterations in a viscous solution are spent converging viscous regions.

To speed up convergence in these regions, each coordinate direction's eigenvalues are multipled by a factor before determining the time step associated with the local spacing and flow conditions. This factor varies from 1.0 when the flow is along the coordinate direction, to the specified value when the flow is perpendicular to the coordinate direction. The far-field CFL number is generally the value specified in the input data file with the CFL# keyword, but near the wall the code effectively increases the time step in the boundary layer by the specified factor. This should increase convergence of the boundary layer.

The default cross-flow CFL factor is 2.0 for three-dimensional flow, and 1.0 (i.e., no time step increase in the boundary layer) for two-dimensional flow and axisymmetric flow.

See Also: CFL#

CYCLES — Number of solution cycles

CYCLES numcyc [PRINT [FREQUENCY] freq]

This keyword is used to set the number of cycles numcyc, and is required in the input data file. By default each cycle has five flowfield iterations in each zone for structured grids, and one for unstructured grids, but this may be changed using the ITERATIONS keyword.

Cycle time information will be written to the .lis file every freq cycles, starting with the first cycle. The default is every cycle.

For structured grids, residuals will also be written to the .lis file every freq cycles, starting with the first cycle, but only for iterations consistent with the PRINT FREQUENCY specified with the ITERATIONS keyword. For unstructured grids, residuals will be written to the <i>.lis</i> file every cycle.

Note that this does not affect integrated properties specified using the LOADS keyword block. The print frequency for integrated properties is controlled by the PRINT keyword in the LOADS keyword block.

Example

If the user specifies

CYCLES 1000 PRINT FREQUENCY 10 ITERATIONS PER CYCLE 10 PRINT FREQUENCY 10

cycle time information will be printed for cycles 1, 11, 21, etc. For structured grids, residuals will only be printed for iterations 10 (in cycle 1), 110 (cycle 11), 210 (cycle 21), etc.

See Also: ITERATIONS, LOADS

${\bf DEBUG-Developer\ debug\ options}$

DEBUG number [mode]

This keyword is intended for use during code development and testing. The various debug options and modes are described in Table 13. For each option, the subroutines referencing that option are listed in parentheses. If *mode* is not described for a given debug option, it should be set to 1 to activate that option. If *mode* is omitted, a default value of 1 is used.

Unless noted otherwise, these options only apply to unstructured grids.

Table 13: Developer Debug Options

number	Description
1	Generate grid and solution files for post-processing by Ensight. (evrzon, US_hdzwrt)
2	Write distance to the nearest wall for each grid point into the .cfl file as the variable WallD. This option applies to both structured and unstructured grids. (InitWallDst, walldfil)
3	Residual computation. (lpschmu, TURB_goldbergUS, TURB_spalartUS, TURB_sstUS)
	$\underline{mode} \underline{ ext{Action}}$
	0 Compute residuals based on $\Delta \mathbf{Q}$
	1 Compute residuals based on right-hand-side
	2 Compute residuals based on right-hand-side normalized by cell volume
4	With the HLLE differencing scheme, replace the anti-diffusive terms with Lax-like dissipation reduction. The size of the dissipation reduction term is scaled by $mode/100$. (hlle, US_HLLE, US_HLLEpt, US_HLLERL)
7	Write the zonal maximum and minimum values of the dependent variables, and the maximum and minimum vertex value of the pressure, into the .lis file. (US_Qcell2Qvrtx)
8	Split the edge data (.cge) file into mode files. (edgopn)
9	Write extra arrays containing cell volume, cell centroid coordinates, face center coordinates, and cell connectivity into the .cge file for debugging purposes. (edgopn, prerdinpt, US_PreUnstructRead, US_PreUnstructWrite)
10	Write detailed grid information and wall distance into the .lis file. For wall distances, this option applies to both structured and unstructured grids. (GlobalWallDist, MkGlblWallLst, US_mklines, US_PreUnstructWrite, US_StencilFill, US_ZnlWallDist)
11	Multiplying factor for viscous-diffusion Jacobian. If DEBUG 11 is not used, a value of 4.0 is used in TURB_goldbergfluxUS, and 2.0*Dissip_Jcb in US_FluxJcbnDiss, where Dissip_Jcb is the Jacobian dissipation parameter set using the LAX DAMPING keyword. (TURB_goldbergfluxUS, US_FluxJcbnDiss)
12	Maximum ratio of production to destruction allowed in Goldberg turbulence model. (TURB_goldbergsrcUS)

Continued on next page

 Table 15: Developer Debug Options (Continued)

number	Description
13	Maximum ratio of destruction to production allowed in Goldberg turbulence model. (TURB_goldbergsrcUS)
14	When updating turbulence model values, write maximum turbulent viscosity to .lis file. (TURB_UpdateGoldberg, TURB_UpdateSA)
15	Not applicable in Wind-US. (NSzsolv, US_unstrpstinp)
16	For multi-species flows, write thermodynamic data table into the file <i>Thermo_Tables</i> . This option applies to both structured and unstructured grids. (chpstin1, propin)
17	When writing flow solution to .cfl file, for vertices at coupled surfaces with multiple boundary conditions (e.g., at corners), make the "coupled" boundary condition the lowest priority. (US_Qcell2Qvrtx)
20	Not applicable in Wind-US. (GRD_UpdateMet)
21	The time step used for the solution of the turbulence model equations is set to <i>mode</i> percent of the time step used for the Navier-Stokes equations. (TURB_lhsdt)
22	Not applicable in Wind-US. (loadqbc, lpmgu, setGridLevel, stagepstinp, US_unstrpstinp)
23	For the Rusanov (Lax) scheme, explicitly set the damping factor for strong shocks to $0.01 \times mode$. The default is to set this damping factor to 0.5 (i.e., equivalent to $mode = 50$). (US_Lax, US_LaxRL)
24	The convergence criterion (see the CONVERGE LOAD keyword) is set to 1./mode. The default convergence criterion is 0.01. (LoadCnvrg)
25	Prevents an error exit when both CFL# AUTOMATIC and a time step in seconds are specified in the same zone. Note though that this shouldn't happen, unless both CFL# AUTOMATIC and CFL MODE 1 or 2 are erroneously used in the same zone. (pstinp)
27	Not applicable in Wind-US. (US_unstrpstinp)
28	Not applicable in Wind-US. (US_unstrpstinp)
30	Dump lines created for line Gauss-Seidel solver to a FieldView "particle path" file for visualization. (US_mklines)
31	Don't stop execution when errors are found in cell volumes or face normals. (US_CheckPreInfo)
32	Minimum line length to be used when creating lines for line Gauss-Seidel solver. The default value is 10. (US_mklines)
35	Use an alternate procedure for computing the friction velocity u_{τ} for wall function boundary conditions. This DEBUG option may only be used for a perfect gas. (US_LawOfTheWall)
38	When implicit boundary conditions are being used (see IMPLICIT BOUNDARY), use explicit boundary conditions at outflow boundaries. (US_rlupw)
39	Compute distance to the nearest wall for each grid point the "old" way (i.e., on a zonal basis). This option applies to both structured and unstructured grids. (MkGlblWallLst, walldfil)

Continued on next page

 Table 15: Developer Debug Options (Continued)

number	Description
41	When implicit boundary conditions are being used (see IMPLICIT BOUNDARY), also treat coupled boundaries implicitly. Requires a parallel run with as many processors as zones, and with direct communication between workers (see the .mpc file communication command in Section 9.6). (US_bcalloc, US_implCpl, US_ImplStencil, US_InitDq, US_NbrCntrbtn, US_rhsvis, US_rlupw, US_TransferDq, wbnd2)
46	Viscous flux correction method.
	$\frac{mode}{0}$ $\frac{\text{Method}}{\text{No correction}}$
	1 Original correction method
	2 Only use normal variation
	3 Only use normal component
	The default is no correction if GRADIENTS LEAST_SQUARES was specified, and the original correction method otherwise. (TURB_diffuseSST, US_FaceConCA, US_FaceConFBnd, US_FaceGradGG)
47	Not applicable in Wind-US. (US_TimeStep)
49	Use full linearization of turbulence model. In addition, if $mode < 0$, and the coefficient matrix is not diagonally dominant, stability will be enhanced by multiplying the relevant diagonal terms by 1.05. (TURB_lhsdt, TURB_sasrcUS)
50	Not applicable in Wind-US. (US_rhsvis)
51	For GRADIENTS LEAST_SQUARES, activate least squares based on face neighbors. (US_LSstencil)
52	When computing viscous fluxes at boundary faces, enforce no shear at reflection boundaries and inviscid walls, and no temperature gradient at viscous adiabatic walls. (US_rhsvisFace)
53	For the Roe scheme, get interface flux using same routines as in structured solver. (This option is currently disabled, and will cause an abort if used in conjunction with RHS ROE.) (US_rhsupw, US_rlupw)
65	Write log of memory allocation/deallocation to the file <i>memdebug.lis</i> . This option applies to both structured and unstructured grids. (MemDebug, prerdinpt, solver_main)

DOWNSTREAM MACH — Outflow boundaries, Mach number

Structured Grids

```
DOWNSTREAM MACH [NUMBER] mach [ZONE] range1[, range2[, ... rangen]]
```

This keyword allows the specification of a mass-averaged subsonic Mach number mach at outflow boundaries in structured grids. Based on the specified mass-averaged Mach number, and using the computed local Mach number and total pressure, a new local static pressure is computed. This boundary condition simulates the uniform Mach number characteristics observed at the compressor faces of turbine engines, and is identical in its effect on the flow to the COMPRESSOR FACE CHUNG keyword.

In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

See Also: COMPRESSOR FACE, DOWNSTREAM PRESSURE, MASS FLOW, TEST 123, TEST 153

DOWNSTREAM PRESSURE — Outflow boundaries, pressure

Structured Grids

```
DOWNSTREAM PRESSURE { \ value [EXTRAPOLATE [SUPERSONIC]] | \ FREESTREAM [EXTRAPOLATE [SUPERSONIC]] | \ EXTRAPOLATE [ALWAYS] | \ value VARIABLE ijk [RELAXER rlxr] | \ value UNSTEADY {SINUSOIDAL | USERSPEC} \Delta p freq phase \ } [ORDER {ZERO|0|0NE|1}] [ZONE] range1[, range2[, . . . rangen]]
```

Unstructured Grids

```
DOWNSTREAM PRESSURE { \ value [EXTRAPOLATE [SUPERSONIC]] | \ FREESTREAM [EXTRAPOLATE [SUPERSONIC]] | \ EXTRAPOLATE [ALWAYS] \ } [ZONE] range1[,range2[, ... rangen]]
```

This keyword controls the specification of a pressure boundary condition at outflow boundaries. In the zone specification, the *range* parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

Structured Grids

There are five possible modes for structured grids, defined as follows:

```
DOWNSTREAM PRESSURE value [EXTRAPOLATE [SUPERSONIC]] \
[ORDER {ZERO|0|ONE|1}] [ZONE] range1[,range2[, ... rangen]]
```

This mode specifies the pressure *value* (in psi) to be applied at the outflow boundary in the specified zone(s). The EXTRAPOLATE [SUPERSONIC] option tells Wind-US to extrapolate all flow conditions to the boundary for points where the flow is supersonic. Either zeroth- or first-order extrapolation will be used, as specified by ORDER.

For boundary points where the flow is subsonic,

- 1. The pressure at the boundary is set to the user-specified value.
- 2. For each boundary point, the density and momentum, plus the effective gamma, compressibility factor, and speed of sound, are extrapolated from the interior to the boundary using either zeroth- or first-order extrapolation, as specified.
- 3. The energy at each boundary point is computed, consistent with the user-specified pressure and the extrapolated values of density, etc.

```
DOWNSTREAM PRESSURE FREESTREAM [EXTRAPOLATE [SUPERSONIC]] \
[ORDER {ZERO|0|ONE|1}] [ZONE] range1[, range2[, ... rangen]]
```

This mode is the same as the previous one, except that the pressure value is taken from the FREESTREAM keyword.

```
DOWNSTREAM PRESSURE EXTRAPOLATE [ALWAYS] \
[ORDER {ZERO|0|ONE|1}] [ZONE] range1[,range2[, ... rangen]]
```

This mode extrapolates all flow conditions for all points on the boundary, regardless of whether or not the flow there is supersonic.

```
DOWNSTREAM PRESSURE value VARIABLE i~j~k [RELAXER rlxr] \ [ORDER {ZERO|0|ONE|1}] [ZONE] range1[, range2[, ... rangen]]
```

This mode imposes the pressure value (in psi) only at the specified (i,j,k) boundary point in the specified zone(s). The indices (i,j,k) should correspond to a point on the outflow boundary, although at present no check is made to verify that they do. The pressure at the remaining boundary points will spatially vary according to the spatial variation of pressure at the solution points adjacent to the boundary, and may be over- or under-relaxed using the input relaxation factor rlxr. The default value for rlxr is 1.0 (i.e., no relaxation).

The procedure is as follows:

- 1. A "reference" pressure p_{ref} at the specified (i, j, k) boundary point is found by extrapolating the interior pressure values to the boundary, using either zeroth- or first-order extrapolation, as specified.
- 2. For each boundary point, the density and momentum, plus the pressure, effective gamma, compressibility factor, and speed of sound, are extrapolated from the interior to the boundary using either zeroth- or first-order extrapolation, as specified.
- 3. At each boundary point, the pressure is re-computed using

$$p = p_{ext} \frac{p_{nom}}{p_{ref}}$$

where p_{ext} is the extrapolated value from step 2, p_{nom} is the nominal value specified with the DOWNSTREAM PRESSURE keyword, and p_{ref} is the "reference" value from step 1.

4. The actual imposed pressure at each point is relaxed using

$$p = rp + (1 - r)p_{old}$$

where r is the input relaxation factor rlxr, and p_{old} is the pressure value from the previous iteration.

5. The energy at each boundary point is computed, consistent with the pressure value from step 4, and the extrapolated values of density, etc., from step 2.

```
DOWNSTREAM PRESSURE value UNSTEADY \ {SINUSOIDAL | USERSPEC} \Delta p freq phase \ [ORDER {ZERO|0|ONE|1}] [ZONE] range1[, range2[, ... rangen]]
```

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This mode allows an unsteady pressure to be applied at the outflow boundary in the specified zone(s). A sinusoidal or user-defined oscillation may be specified. The parameters are defined as:

value Baseline pressure, p_s (psi) Δp Amplitude of oscillation (psi) freq Frequency of oscillation, ω (Hz) phase Phase angle of oscillation, ϕ (deg)

If SINUSOIDAL is used, specifying a sinusoidal pressure oscillation, the pressure is computed from

$$p = p_s + \Delta p \sin[2\pi(\omega t + \phi)]$$

where p is the pressure and t is the time.

If USERSPEC is used, specifying a user-defined pressure oscillation, a table must be input immediately following the DOWNSTREAM PRESSURE keyword, with the following format:

```
PROFILE npts

per(1) amp(1)

per(2) amp(2)

...

per(npts) amp(npts)
```

where npts is the number of points in the profile (max 65), per is the normalized period (i.e., from 0.0 to 1.0), and amp is the normalized amplitude (i.e., from -1.0 to 1.0). As an example, the following normalized pressure oscillation:

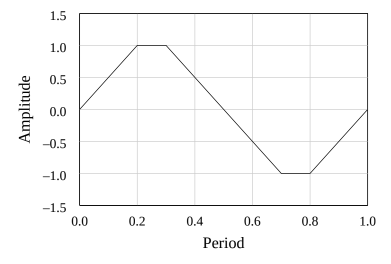


Figure 7: Example user-specified unsteady pressure oscillation

could be specified using the following profile input:

PROFILE 6 0.0 0.0 0.2 1.0 0.3 1.0 0.7 -1.0

```
0.8 -1.0
1.0 0.0
```

Once the downstream pressure is determined for the current time level, the density and momentum at each boundary point, plus the effective gamma, compressibility factor, and speed of sound, are extrapolated from the interior to the boundary using either zeroth- or first-order extrapolation, as specified. The energy at each boundary point is then computed, consistent with the pressure and the extrapolated values of density, etc.

Extrapolation Notes

The default for all extrapolation is zeroth-order (i.e., conditions at the boundary are set to the values at the computational plane adjacent to the boundary). This results in a discontinuous slope in flow values near the outflow boundary, which may be important for flows with significant streamwise pressure gradients. First-order extrapolation yields smoother results.

For flows with little or no streamwise pressure gradient near the outflow boundary, the results using zeroth- and first-order extrapolation are essentially identical. Convergence rates and the final residual values are generally better with zeroth-order extrapolation, however, so the default zeroth-order extrapolation is recommended.

For flows with significant streamwise pressure gradients near outflow boundaries, zeroth-order extrapolation can give poor results at the outflow boundary, and in some cases these can affect values at the inflow boundary. First-order extrapolation is thus recommended for these flows.

```
See Also: COMPRESSOR FACE, DOWNSTREAM MACH, MASS FLOW, FREESTREAM, TEST 123, TEST 153
```

Unstructured Grids

There are three possible modes for unstructured grids, defined as follows:

```
DOWNSTREAM PRESSURE value [EXTRAPOLATE [SUPERSONIC]] \ [ZONE] range1 [, range2 [, . . . rangen]]
```

This mode specifies the pressure *value* to be applied at the outflow boundary in the specified zone(s). The EXTRAPOLATE [SUPERSONIC] option tells Wind-US to extrapolate all flow conditions to the boundary for points where the flow is supersonic. For unstructured grids, only zeroth-order extrapolation is available.

```
DOWNSTREAM PRESSURE FREESTREAM [EXTRAPOLATE [SUPERSONIC]] \
[ZONE] range1[,range2[, ... rangen]]
```

This mode is the same as the previous one, except that the pressure value is taken from the FREESTREAM keyword.

```
DOWNSTREAM PRESSURE EXTRAPOLATE [ALWAYS] \
[ZONE] range1[,range2[, ... rangen]]
```

This mode extrapolates all flow conditions for all points on the boundary, regardless of whether or not the flow there is supersonic.

```
See Also: MASS FLOW, FREESTREAM, TEST 123, TEST 153
```

$DQ - \Delta Q$ limiter

```
DQ [LIMITER] [OFF | [ON | NXAIR] [DRMAX drmax] [DTMAX dtmax] \ [DPMAX dplim] [RELAX dqrlx]]
```

The DQ LIMITER keyword may be used to limit the change in the solution over a single iteration. This may be helpful, especially in the initial stages of a calculation, in maintaining stability, and allow the use of larger CFL numbers than might otherwise be possible.

If the DQ LIMITER keyword is not used:

- If all zones are structured, the $\Delta \mathbf{Q}$ limiter is off.
- If all zones are unstructured, the $\Delta \mathbf{Q}$ limiter is on.
- If there is a mix of structured and unstructured zones, and CFL# AUTOMATIC is not used in any zone, the $\Delta \mathbf{Q}$ limiter is off.
- If there is a mix of structured and unstructured zones, and CFL# AUTOMATIC is used in any zone, the $\Delta \mathbf{Q}$ limiter is on.

Note that although the DQ LIMITER keyword has separate options for structured and unstructured zones, it is not zonal-dependent. The limiter is either off in all zones (both structured and unstructured), or on in all zones using the default or user-specified limits.

Structured Grids

For structured grid zones, the keyword limits the solution by specifying limits on the change in density and/or temperature over a single iteration. The following options apply to structured grid zones.

OFF	Don't limit the solution.
ON	Use the MacCormack $\Delta \mathbf{Q}$ limiter (MacCormack, 1997). With this limiter, the entire solution, including the turbulence variables, is limited throughout the zone, such that the specified fractional changes in density and temperature are not exceeded. (See below for some further considerations when multiple subiterations are being used.)
NXAIR	Use the $\Delta \mathbf{Q}$ limiter from the NXAIR code. This limiter acts locally; the solution is limited only at those points that exceed the specified fractional changes in density or temperature. Unlike the MacCormack limiter, the NXAIR limiter is not applied to the turbulence model equations (although the models of course use the limited Navier-Stokes variables, so there is an indirect influence).
	The NXAIR limiter may only be used with perfect gas flows.
DRMAX $drmax$	drmax specifies the fractional change allowed in the density over an iteration. I.e., specifying "DQ LIMITER ON DRMAX 0.2" means the density will not be permitted to change by more than $\pm 20\%$ from one iteration to the next. The default value is 0.5.
$\mathtt{DTMAX}\ dtmax$	dtmax specifies the fractional change allowed in the temperature over an itera-

tion. The default value is 0.5.

If just DQ LIMITER is specified, without specifying any options, the MacCormack $\Delta \mathbf{Q}$ limiter is used. Since the defaults for drmax and dtmax are both 0.5. both density and temperature will be allowed to change by no more than $\pm 50\%$ over an iteration.

The use of the DQ LIMITER ON keyword becomes somewhat problematic when multiple subiterations are used for the Navier-Stokes equations (using the NAVIER-STOKES ITERATIONS keyword) and/or the turbulence model equations (using the ITERATIONS option with the TURBULENCE keyword). By default, the number of sub-iterations for both the Navier-Stokes and turbulence model equations is one. In this case, each iteration for the Navier-Stokes equations is followed by an iteration for the turbulence model equations, and if the limiter is applied to the Navier-Stokes solution, it is also applied to the turbulence model solution.

If the number of Navier-Stokes sub-iterations is one, and if multiple sub-iterations are used for the turbulence model equations, when the limiter is applied to the Navier-Stokes solution it is applied to each turbulence model sub-iteration.

Also, if multiple Navier-Stokes sub-iterations are used, it's possible that the computed flow field is such that the limiter is applied during some sub-iterations, but not others. In this case, if the limiter is applied during the final Navier-Stokes sub-iteration, it is again applied to each turbulence model sub-iteration. If the limiter is not applied during the final Navier-Stokes sub-iteration, it will not be applied at all during the turbulence model iteration.

For these reasons, it is recommended that the number of sub-iterations for the Navier-Stokes and turbulence model equations be left at their default values of one when the DQ LIMITER ON keyword is used.

Unstructured Grids

For unstructured grid zones, the keyword limits the solution by specifying a relaxation factor, and a limit on the change in pressure over a single iteration. The following options apply to unstructured grid zones.

OFF Don't limit the solution.

ON Apply the $\Delta \mathbf{Q}$ limiter using the default or user-specified limits.

DPMAX dplim dplim specifies the maximum allowable change in pressure, as a fraction of the

local pressure, over an iteration. I.e., specifying "DQ LIMITER ON DPMAX 0.2" means the solution will be limited such that the pressure doesn't change by more

than 20% from one iteration to the next. The default is 0.1.

RELAX dqrlx Relaxation factor. The default is 1.0 if a first-order explicit operator (specified

by the RHS keyword) is being used in all zones, and 0.5 otherwise.

If the DQ LIMITER keyword is used, but without specifying OFF or ON, the ΔQ limiter is on.

When the $\Delta \mathbf{Q}$ limiter is off, no solution limiting is done. When it's on, the entire solution, including the turbulence solution, is always limited using the relaxation factor dqrlx. In addition, the Navier-Stokes and chemistry solution is limited such that the density changes by no more than 50% over an iteration, and the pressure changes by no more than the amount specified using the DQ LIMITER keyword. And the turbulence solution is limited such that the turbulence variables change by no more than 50% over an iteration.

DQ LIMITER OFF may not be specified if CFL# AUTOMATIC is used in any zone.

See Also: Q LIMIT

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${\bf END-Termination}$

[END]

This keyword, which must begin in column 1, may be used to end the input data file.

${\bf ENTHALPY\text{-}PRESERVING--Enthalpy\text{-}preserving scheme}$

Unstructured Grids

ENTHALPY-PRESERVING

When this keyword is used, the left and right states used to compute the cell interface fluxes are modified based on an enthalpy-preserving scheme. The default is to not use an enthalpy-preserving scheme.

ENTROPY FIX — Entropy fix for Roe scheme

ENTROPY [FIX] [ON|OFF]

When the Roe scheme is being used, this keyword may be used to turn off a partial "entropy fix", intended to help prevent expansion shocks by not allowing the flow eigenvalues to have non-physical values. The default is ON.

See Also: RHS

EXTRAPOLATE — Extrapolation at freestream, arbitrary inflow boundaries

EXTRAPOLATE [BOUNDARY] [ORDER] ZERO | O | ONE | 1 [zone_selector]

This keyword sets the extrapolation order used for boundary conditions at freestream and arbitrary inflow boundaries. The default is zeroth-order extrapolation which should be more stable. However, for grids with good mesh resolution near the boundary, first-order is recommended, and should be more accurate.

 $See\ Also:$ ARBITRARY INFLOW

FIXED_CL — Fixed C_L computation

This keyword may be used to vary the angle of attack to obtain a specified lift coefficient for surface(s) defined by the SURFACE and/or SUBSET keywords (with FORCE specified) in the LOADS keyword block. Note that the specified lift coefficient is the total for all surfaces combined.

cltarg The desired lift coefficient. The default is 0.0.

dalim Limit in the angle of attack change allowed each update (degrees). The default is 0.2

degrees.

rlxalph Relaxation parameter. The default is 1.0, for no relaxation.

 $\it alfafreq$ Update frequency, in cycles. The default is 1, to update every cycle.

istart Starting cycle. The default is 1, to start on the first cycle.

By default, the angle of attack changes are written to the .lis file. Specifying HISTORY OFF will prevent this.

See Also: LOADS, TEST 11

FIXER — Instability smoothing

Structured Grids

```
FIXER [PRINT] [zone_selector]
```

Unstructured Grids

```
FIXER [FIRST ORDER] [MAX max\_pts AFTER iter] [zone\_selector]
```

Wind-US's fixer capability searches for computational points at which numerical instabilities have resulted in physically impossible results, and replaces the faulty values with averages from nearby points. Using fixer may help the solution through difficult transients, but it may also simply smear out instabilities over a larger portion of the flowfield. You should maintain a careful watch over the solution.

Structured Grids

For structured grids, values at points where the density or internal energy is negative are replaced with averages from nearby j and k points in the same i-plane. (I.e. the smoothing is two-dimensional; values from adjacent i-planes are not used.) The process is iterative, with multiple smoothing passes. Only "active" points are examined, not wall, coupled boundary, hole, or fringe points.

In a given i-plane, if more than 50% of the points are "bad", or if the smoothing process fails to remove all the negative values, the run is aborted.

For each i-plane containing negative values of density or internal energy, the zone, iteration number, i index, and number of bad points will be written to the list output (.lis) file. If the PRINT option is specified, the listing will also include the j and k indices of each bad point.

Unstructured Grids

Values at points where the internal energy, density, or pressure is outside the allowed limits are replaced with averages from neighboring cells. The limits on density and pressure may be specified using the $\tt Q$ LIMIT keyword, and the limits on internal energy are computed from those. The defaults, normalized by the freestream static values, are 0.01 and 250 for both density and pressure.³²

If FIRST ORDER is specified, the bad cells will be flagged and a first order scheme will be used for them for the remainder of the computation.

The MAX ... keyword may be added to specify that the maximum number of bad cells allowed is max_pts , but don't start enforcing this until iteration iter. If the limit is exceeded, the run is aborted. If MAX ... is not used, no limit is set on the maximum number of bad cells.

See Also: Q LIMIT

 $^{^{32}}$ For frozen and finite-rate chemistry, only density and pressure are checked.

${\bf FREESTREAM--Freestream\ conditions}$

FREESTREAM {STATIC | TOTAL} [CONDITIONS] $M~P~T~\alpha~\beta$

This keyword allows input of freestream flow conditions. The user-specified parameters are defined as:

M Mach number

P Static or total pressure (psi), depending on the STATIC | TOTAL keyword

T Static or total temperature (°R), depending on the STATIC | TOTAL keyword

 α , β Angles of attack and yaw (degrees)

Note that, if U is the freestream velocity magnitude, α and β are related to the Cartesian velocity components by:

$$w = U \sin \beta$$
$$u = U \cos \beta \cos \alpha$$
$$v = U \cos \beta \sin \alpha$$

Note: The local flow angles (localpha and locbeta in the CFPOST post-processing package) are defined relative to the x, y, z coordinates and are given by:

$$localpha = tan^{-1}(v/u)$$

 $locbeta = tan^{-1}(w/u)$

Thus if β is not equal to 0, *locbeta* is not equal to β .

Note: Wind-US assumes α is rotation about the z axis. That is, y is up.

See Also: INITIALIZE, REINITIALIZE

FRINGE — Solution mode at fringe points

Structured Grids

FRINGE [MODE] FROZEN [$zone_selector$]

For overlapping structured zonal boundaries, this keyword may be used to freeze the solution at fringe points during a cycle. Solution values at these points are then updated at the end of the cycle.

FRINGE FROZEN must be used for double-fringe boundaries. Running a double-fringe grid without this keyword may lead to erroneous results.

FRINGE FROZEN may also be specified for single-fringe boundaries, but is not required. The default for single-fringe boundaries is to update the solution at fringe points every iteration.

Using double fringes increases the solution accuracy at overlapping boundaries. Note, though, that by freezing the solution over a cycle, the solution at fringe points will lag in time behind the solution at interior points. This could potentially lead to instabilities during transients, or a reduction in the convergence rate if a large number of iterations per cycle are specified. However, testing on a limited number of cases has not revealed any problems when using up to five iterations per cycle.

Double-fringe boundaries may be created with GMAN, by first selecting double-fringe mode using (in GMAN's graphical mode) MODIFY BNDY > GENERATE FRINGE > SET FRINGE-MOD > DOUBLE FRINGE, then generating the fringe.

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GAS — Gas property specification

${\tt GAS} \ gam \ prl \ prt \ gasc$

This option allows the user to input different values of γ , laminar and turbulent Prandtl numbers, and gas constant. The parameters are defined as:

gam Ratio of specific heats $\gamma = c_p/c_v$ (default is 1.4) prl Laminar Prandtl number (default is 0.72) prt Turbulent Prandtl number (default is 0.90) gasc Gas constant R (default is 1716 ft²/sec²-°R)

Note: The ARBITRARY INFLOW keyword block, if used, must come after the GAS keyword in the input data (.dat) file.

GRADIENTS — Gradient computation

Unstructured Grids

GRADIENTS {CELL_AVERAGE | LEAST_SQUARES} [zone_selector]

The default gradient computation procedure in Wind-US for unstructured grids (CELL_AVERAGE) is a carry-over from the initial unstructured solver in Wind-US 1.0. Average face values are first computed using an inverse distance weighted average among the cell-centered values from cells sharing the face. The Green-Gauss theorem is then used to compute the average gradient in the cell from the average face values. In the unstructured solver introduced in the current version of Wind-US, this has been shown to be stable, but can be inaccurate, especially for high-aspect-ratio cells, and is not guaranteed to preserve an initially-linear arbitrary function in the computational mesh.

Specifying LEAST_SQUARES causes gradients to be computed using a least squares fit to the flow field values in the region of the cell. Inverse distance weighting is used, with values obtained from neighboring faces or nodes (Mavriplis, 2003). This procedure should be specified whenever second-order accuracy is important, especially with highly-stretched meshes typically used for viscous problems. For example, for computing accurate drag values on a body at cruise conditions, using LEAST_SQUARES is essential.

LEAST_SQUARES may only be used for perfect gases.

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${\bf GRAVITY-Add\ gravity\ body\ forces}$

GRAVITY x_c y_c z_c g_x g_y g_z

The GRAVITY keyword allows the user to include the effects of gravity on the fluid.

- x_c, y_c, z_c Coordinates of a point on the "ground" plane (normal to the gravity vector) used to define the location of zero potential energy, in grid units
- g_x , g_y , g_z Components of the gravity vector, in ft/sec². For example, if the y direction is normal to the earth's surface, these values might be 0.0, -32.2, 0.0.

This keyword cannot be used with the Van Leer differencing scheme (RHS VANLEER).

GRID LIMITER — Grid limiting capability

GRID LIMITER {ANGLE $angle \mid OFF$ } [$zone_selector$]

Wind-US's grid limiting capability switches the explicit operator to a first-order scheme in the presence of grid turning greater than a specified amount. This capability may be needed to reduce numerical instabilities near wing tips, leading edges, and other geometric features which may require extreme amounts of grid turning.

The parameter *angle* is the number of degrees off of a straight line allowed before limiting the scheme to first order.

This keyword applies to most of the first-, second-, and third-order explicit operators available with the RHS keyword. It does not apply to the CENTRAL and COAKLEY schemes (but see TEST 139 for COAKLEY). It also does not apply to first-order upwind schemes, and may not be used with third-order fully upwind, fourth-order, or fifth-order schemes.

See Also: TEST 110, TEST 139, RHS, TVD

HEAT-EXCHANGER — Heat exchanger model model (block)

Structured Grids

```
HEAT-EXCHANGER

ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \

[SUBSET I range J range K range]

ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \

[SUBSET I range J range K range]

TEMPERATURE [GAIN] dtemp

CHX cploss EXPONENT exp

END HEAT-EXCHANGER
```

This keyword block enables the user to model the effects of heat exchangers by specifying a discontinuous change in properties across a zone boundary or portion of a zone boundary. Multiple heat exchangers may be modeled by using multiple HEAT-EXCHANGER keyword blocks The following restrictions apply:

- This keyword applies to structured grids only.
- The BOUNDARY TVD FACTOR 0 keyword option should be used for all zonal boundaries with heat exchangers.

The various elements of the HEAT-EXCHANGER input block are defined as follows:

HEAT-EXCHANGER

Defines the beginning of the heat exchanger keyword input block.

```
END HEAT-EXCHANGER
```

Ends the heat exchanger input block

```
ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \ [SUBSET I range J range K range]

ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \ [SUBSET I range J range K range]
```

These two lines define the location of the heat exchanger. The relevant zones are given by the values of iz1 and iz2, and the relevant boundaries within zones iz1 and iz2 are specified via the BOUNDARY keyword parameter.

- iz1 First zone containing heat exchanger (i.e., the zone coupled to zone iz2).
- iz2 Second zone containing heat exchanger (i.e., the zone coupled to zone iz1).

At the specified boundaries in the above zones, if the flow is leaving the zone the heat exchanger will be modeled by increasing the temperature and decreasing the pressure when passing flow conditions to the other (i.e., downstream) zone. Conversely, if the flow is entering the zone, the temperature will be decreased and the pressure increased when passing flow conditions back to the upstream zone.

The SUBSET parameter may be used to specify that the change in secondary velocity occurs only over a part of the zone boundary. Otherwise, it is assumed that the change occurs over the entire boundary. The *range* parameters define the part of the zone boundary over which the change occurs, and take one of the following forms:

index1 index2 Starting and ending indices in the specified direction. LAST may be used for the last index.

ALL Equivalent to 1 LAST.

The starting and ending indices for the appropriate I, J, or K parameter (depending on the boundary specified) must be the same, and correspond to that boundary.

TEMPERATURE [GAIN] dtemp

This defines the temperature change across heat exchanger, dtemp, in °R.

 ${\tt CHX} \ cploss \ {\tt EXPONENT} \ exp$

This defines the loss across the heat exchanger. The input value *cploss* is the static pressure loss coefficient, $\Delta p/(\rho u)^n$, and exp is the mass flow rate exponent, n.

HISTORY — Time history flowfield variable tracking (block)

Structured Grids

```
HISTORY

VARIABLE [M] [p] [T] [u] [v] [w] [radius] [DeltaP] [x] [y] [z] \

[rho] [rho*u] [rho*v] ...

[FREQUENCY f]

REGION nzn ibeg iend jbeg jend kbeg kend

REGION nzn {NODE | CELL} num

...

ENDHISTORY ...
```

For unsteady flow problems, the HISTORY keyword block allows the user to track the temporal evolution of variables at discrete specified points. The specified flow quantities are written to a common time history (.cth) file every f iterations. Upon completion of a time history run, the auxiliary program thplt may be used to analyze the data and/or to create a GENPLOT file that may then be plotted using CFPOST.³³ The .cth file contains reference and scaling data. Thus, when thplt is used to create a GENPLOT file, the time history variables will be dimensional.

For the results to be strictly valid, all zones must run with the same time step and with one iteration per cycle. However, in recognition that this is not always feasible, this capability can be used with only the stipulation that the zone(s) in which temporal evolution information is desired be run with a constant time step.

The .cth file itself does not support direct I/O. Therefore, when Wind-US is being run in parallel mode, each individual processor maintains the requested time history for its zones, but calls the master process to update the .cth file whenever the flow (.cfl) file is updated, and whenever the time history memory buffer is filled. This forces the .cth file to be consistent with the .cfl file.

It should be noted that although time history files are commonly referred to as .cth files, Wind-US automatically appends the current date and time to the end of the file name, using the format mm-dd-yyhhhmmm. As an example, for a case run using an input data (.dat) file named config3.dat, the resulting time history file might be config3.cth.08-22-07h22m19, indicating that it was created at on Aug. 22, 2007, at 10:19 PM. Restarting a calculation will thus result in multiple time history files, one for each individual run. The time history post-processing tool thplt can read multiple .cth files to analyze the data and/or create a single GENPLOT file combining the different time intervals.

The various elements of the HISTORY keyword block are defined as follows:

HISTORY

Defines the beginning of the HISTORY block

Specifies the variables to be recorded. At least one must be specified, and the maximum allowed is fifteen. Note that variable names are *case-sensitive*. The first eight variables shown above are defined as follows:

 $^{^{33}}$ Because of changes to the format of the .cth file, the timplt utility used to analyze .cth files created by WIND 5.51 and earlier will not work with .cth files created by the current Wind-US code.

M	Mach number
p	Static pressure
T	Static temperature
u	Velocity in the x direction
v	Velocity in the y direction
W	Velocity in the z direction
radius	Distance from a specified reference point to the location of a shock front. See
	the section Front Tracking Mode for details.
DeltaP	Monotone pressure rise across the shock front. See the section Front Tracking
	Mode for details.

In addition to the variables listed above, any variable in the q array may be specified. These of course depend on the equations being solved. The possible q variables are listed in Table 14, in groups, along with an indication of when they are available.

Table 14: HISTORY Keyword — q Variables

Name	Definition When Used		
—Navier-Stokes equations			
rho	Static density	Always	
rho*u	Momentum in the x direction		
rho*v	Momentum in the y direction		
rho*w	Momentum in the z direction		
rho*e0	Total energy		
—Auxiliary parameters			
beta	Effective specific heat ratio	Always	
Z	Compressibility factor		
a	Speed of sound		
kappa	Thermal conductivity		
dt	Time step		
Viscosity			
mul	Laminar viscosity coefficient	Viscous flows	
mut	Turbulent viscosity coefficient		
— Turbulence	e equations		
anut	Eddy viscosity	Baldwin-Barth and	
		Spalart-Allmaras turbulence models	
k	Turbulent kinetic energy	SST and Chien k - ϵ	
		turbulence models	
omega	Specific dissipation rate	SST turbulence model	
epsilon	Turbulent dissipation	Chien k - ϵ turbulence model	

Continued on next page

Table 14: HISTORY Keyword — q Variables (Continued)

Name Definition		When Used		
—Chemistry equations				
species	Mass fraction of <i>species</i> , where <i>species</i> is the species name defined in the chemistry data file (see Section 7.11 and the CHEMISTRY keyword block)	Non-reacting chemistry		
${\tt rho} {\star} species$		Finite-rate reacting chemistry		
shf	Specific heat of formation	Chemistry		
—Navier-Stokes equations in rotating frame				
rho*ur	Momentum in the x direction in a rotating reference frame (rho*u, etc., are in the inertial frame)	Rotating reference frame		
rho*vr	Momentum in the y direction in a rotating reference frame			
rho*wr	Momentum in the z direction in a rotating reference frame			
rho*e0r	Total energy in a rotating reference frame			
-MFD equat	tions			
Bx, By, Bz	Magnetic field components in the $x, y,$ and z directions	MFD flows		
Ex, Ey, Ez	Electric field components in the $x, y,$ and z directions			
Jx, Jy, Jz	Current density components in the x , y , and z directions			
Lx, Ly, Lz	Lorentz force components in the $x, y,$ and z directions			
sigma	Conductivity			
voltage	Voltage			

FREQUENCY f

Specifies the number of solver iterations between output of time history information. The default is to write to the .cth file every 10 iterations.

 ${\tt REGION} \ nzn \ ibeg \ iend \ jbeg \ jend \ kbeg \ kend$ REGION nzn {NODE | CELL} num. . .

Defines the discrete locations where temporal information will be sampled in zone nzn. The different formats correspond to a structured zone range, and a particular node or cell in an unstructured node-based or cell-based scheme.³⁴

³⁴Note, though, that Wind-US does not yet support the use of the HISTORY keyword with unstructured grids.

Except for the special-purpose variables radius and DeltaP, in a structured grid time history data will be recorded at each point in the domain defined by the indices (*ibeg,jbeg,kbeg*) to (*iend,jend,kend*). The variables radius and DeltaP are intended for use in tracking the location of a shock wave. In this case, the REGION parameters are defined differently. See the section Front Tracking Mode for details.

Multiple REGION lines may be used to specify multiple sampling regions, in the same zone, or different zones. Grid sequencing is supported for structured grids, but there is presently no other way (except multiple REGION lines) to spatially sub-sample a region.

ENDHISTORY

Defines the end of the HISTORY block.

Front Tracking Mode

For structured grids, a special mode exists that can be useful for tracking the motion of shock waves. If the region is specified as:

```
REGION nzn i0 idir j0 jdir k0 kdir
```

where one of idir, jdir, kdir is ± 1 and the other two are zero, then front tracking mode is active in zone nzn.

Beginning at the point $(i\theta,j\theta,k\theta)$, the solution is sampled to the end of the zone in the specified direction, looking for the maximum change in pressure across a cell. This location is then used as the "probe location" to output the specified variables. In this context, the radius variable represents the distance of this point from the point $(i\theta,j\theta,k\theta)$. Note that this is probably the only time that a time history variable of x, y, or z makes sense, as they can be used to more precisely specify the location of the detected front.

To provide an indication of the evolution of the shock strength, the variable DeltaP can be used. When this variable is specified, the search is continued from the detected front location to the end of the zone, in the specified search direction, and the maximum monotone pressure change is recorded.

Example 1

The following use of the HISTORY block will create a time history file containing the static pressure at the points (17,1,1), (17,11,5), and (9,11,7) in zones 1, 2, and 3, respectively. Values will be written to the file every ten iterations.

```
History
Variable p
Region 1 17 17 1 1 1 1 1
Region 2 17 17 11 11 5 5
Region 3 9 9 11 11 7 7
Endhistory
```

Example 2

In the following example, the Mach number and pressure will be recorded in zone 5, for the grid points (i, j, k) = (7-16,22-24,19). Values will be written to the file every five iterations.

```
History
Variable M p
Frequency 5
```

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$Example \ 3$

In this example, the position and strength of a shock wave will be tracked in the +i direction in zone 4. The point (1,52,20) will be used as the reference point, and values will be recorded every 20 iterations.

History
Variable radius x DeltaP
Frequency 20
Region 4 1 1 52 0 20 0
Endhistory

HLLE | RUSANOV — Anti-diffusion terms for HLLE and Rusanov schemes

Structured Grids

 $\{ \verb+HLLE | \verb+RUSANOV+ EPSS + epss [zone_selector] \\$

This keyword allows the user to control the behavior of the anti-diffusion term in the HLLE and Rusanov schemes for structured grids. The parameter epss is used in the computation of a switch used to regulate the anti-diffusion terms. The possible values for epss are shown below.

epss	Switch	Result
0.0	1.0	Anti-diffusion terms are always fully on.
> 0.0	$1.0 - epss p_{scal} $	Anti-diffusion terms are on, scaled by the absolute value of a pressure gradient scaling parameter p_{scal} , with $epss$ as a coefficient. The result is that the anti-diffusion terms are smaller in regions where the pressure gradient is changing rapidly. Larger values of $epss$ result in smaller anti-diffusion terms. The switch is limited to a minimum value of 0.0 (fully off).
< 0.0	1.0 + epss	Anti-diffusion terms are on, with the switch hard-wired to the indicated value. Larger negative values of $epss$ result in smaller anti-diffusion terms. The switch is limited to a minimum value of 0.0 (fully off), which corresponds to $epss = -1.0$.

The default value for epss is 0.0 for the HLLE scheme, and 0.5 for the Rusanov scheme.

See Also: RHS

HOLD — Hold conditions at freestream inflow boundaries

HOLD {TOTALS | CHARACTERISTICS} [zone_selector]

Specifying HOLD TOTALS indicates that total conditions are to be held constant at freestream boundaries with inflow. For unstructured grids, this is only valid for an ideal gas. Specifying HOLD CHARACTERISTICS indicates that characteristic values are to be held constant. Note that the total pressure is always held fixed at freestream inflow boundaries, whether HOLD TOTALS is specified or not. The option specified will be applied at all the freestream boundaries in the specified zone(s).

The default is HOLD CHARACTERISTICS.

Note that the HOLD keyword only applies to freestream boundaries with subsonic inflow. See the HOLD_TOTALS keyword in the ARBITRARY INFLOW keyword block for information on holding total conditions at arbitrary inflow boundaries.

Note also that the syntax is slightly different for freestream and arbitrary inflow boundaries. For freestream boundaries, HOLD TOTALS and HOLD CHARACTERISTICS are used, without an underscore. For arbitrary inflow boundaries, HOLD_TOTALS and HOLD_CHARACTERISTICS are used in the ARBITRARY INFLOW keyword block, with an underscore.

IMPLICIT — Implicit operator control

Structured Grids

Unstructured Grids

This keyword allows user control of the implicit operator within each zone. There are five possible modes for structured grids, and two for unstructured grids.

Structured Grids

```
IMPLICIT word1 [word2 word3 [zone_selector [tre]]]
```

This form of the IMPLICIT keyword allows the user to turn off the implicit operator completely, resulting in an explicit calculation, or to use a scalar or full block implicit operator. With these options, a different implicit operator may be specified for each computational direction.

The user-specified parameters are defined as follows:

```
      word1
      A keyword controlling the \xi-direction implicit operator

      word2
      A keyword controlling the \eta-direction implicit operator

      word3
      A keyword controlling the \zeta-direction implicit operator

      tre
      Specifies the trapezoidal time differencing factor. The possible values are:

      0
      Explicit (but expensive, don't use)

      1 or omitted
      Implicit

      0.5
      Trapezoidal time differencing
```

The keywords that may be used for word1, word2, and word3 are defined in the following table.

Keyword	Difference Scheme
NONE	Explicit
SCALAR	Scalar implicit operator (default for Euler solutions)
FULL	Full block implicit operator (default for viscous solutions)

Note that if only one of the above keywords is specified, Wind-US assumes that the specified operator is to be used in all directions in all zones.

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```
IMPLICIT JACOBI jiter jcnv [RELAX jrel] [zone_selector]
```

This form of the IMPLICIT keyword turns on the point Jacobi implicit operator in the selected zone. The user-specified parameters are defined as follows:

The number of subiterations allowed each time step. A typical value is 30.

icnvThe level of convergence to assume the subiterations are converged. A typical value is 0.0001.

jrelThe relaxation factor. The default value is 1.0 (i.e., no relaxation).

```
IMPLICIT GAUSS giter gcnv [RELAX grel] [zone_selector]
```

This form of the IMPLICIT keyword turns on the Gauss-Seidel implicit operator in the selected zone. The user-specified parameters are defined as follows:

qiterThe number of subiterations allowed each time step. A typical value is 10.

qcnvThe level of convergence to assume the subiterations are converged. A typical value is 0.0001.

The relaxation factor. The default value is 1.0 (i.e., no relaxation). qrel

```
IMPLICIT MACCORMACK [MAFK k] [zone_selector]
```

This form of the IMPLICIT keyword turns on MacCormack's first-order modified approximate factorization (MAFk) procedure in the selected zone. The procedure attempts to remove the decomposition error by feeding the error term back into the matrix equations on the right-hand side.

The solution process requires subiterations, with the number of subiterations specified with the MAFK option by the parameter k. The default for k is 2.

For stability, the IMPLICIT BOUNDARY keyword should be used to specify that implicit boundary conditions are to be used with the MAFk procedure. In addition, when large CFL numbers are to be used, it is recommended that the CFL number be increased gradually (over 200 iterations or so) to the desired value using the INCREMENT parameter in the CFL# keyword.

The MAFk procedure has been demonstrated in Wind-US to be stable in two dimensions with very high CFL numbers (greater than 1000). In three dimensions, however, only limited testing has been done, and its efficiency has not yet been determined.

The MAFk procedure may not be used with second-order time marching.

```
IMPLICIT OVERFLOW [zone_selector]
```

This form of the IMPLICIT keyword specifies that the "ARC3D 3-factor diagonal scheme" as implemented in OVERFLOW 1.8q is to be used. Tests indicate that this scheme is faster than the other implicit schemes, and gives comparable answers.

This option is currently available only for 3-d perfect gas flows, with explicit boundary conditions. It also uses more memory than the other implicit schemes, due to the interface coding used to implement it in Wind-US.

See Also: IMPLICIT BOUNDARY, IMPLICIT ORDER

Unstructured Grids

IMPLICIT NONE [zone_selector]

This form of the IMPLICIT keyword allows the user to turn off the implicit operator completely, resulting in an explicit calculation.

This form of the IMPLICIT keyword a specifies that a Gauss-Seidel implicit operator is to be used in the selected zone. The various options are as follows:

BLOCK | DIAGONAL Specifies that the equations are to be solved as either a full block or

diagonal matrix system. The default is a full block matrix system.

LINE Solve the Navier-Stokes equations using a line Gauss-Seidel implicit

solver. The default is a point Gauss-Seidel implicit solver. If this option

is used, STORE_EXACT_LHS must also be specified.

Note that even when LINE is specified, the turbulence equations will be

solved using a point Gauss-Seidel solver.

VISCOUS_JACOBIAN {FULL|SCALAR}

Use either a full or scalar (diagonal approximate) viscous Jacobian. The

default is SCALAR.

SUBITERATIONS nsub Specifies the number of sub-iterations to be performed on each face, for

each "iteration per cycle," for the point Gauss-Seidel solver. The default

is 10.

STORE_EXACT_LHS Store the entire left-hand-side in memory. The default is a low-storage

implicit algorithm with an approximate Jacobian based on the Rusanov

scheme.

CONVERGE [ORDER order] [FREQUENCY freq]

Specifies the convergence criteria for the sub-iterations for the point Gauss-Seidel solver (i.e., the LINE option is not being used). order is the desired order of convergence, and freq is the frequency for checking convergence, in terms of Gauss-Seidel sub-iterations (i.e., the value specified using SUBITERATIONS). This only applies when STORE_EXACT_LHS is specified, and CFL# AUTOMATIC is being used. The defaults for order

and freq are 2 and 5, respectively.

The following additional options specify actions that are now always done in the current version of Wind-US. They have been retained for backward compatibility, so that existing input data (.dat) files may be used, but otherwise have no effect.

JACOBIAN [PER] FACE Specifies that the flux Jacobian is to be computed on each face.

SAVE_JACOBIAN Specifies that the Jacobians are to be saved between iterations.

IMPLICIT BOUNDARY — Implicit boundary conditions

IMPLICIT BOUNDARY {ON | $\overline{\text{OFF}}$ } [$zone_selector$]

This keyword controls the use of Wind-US's numerical boundary conditions within the implicit operator. The default mode is OFF, implying explicit boundary conditions. When ON is specified, in structured grids implicit boundary conditions will be used at reflection boundaries, and viscous and inviscid walls (with and without bleed or blowing). In unstructured grids, implicit boundary conditions will be used at reflection boundaries, viscous and inviscid walls (without bleed or blowing), and outflow boundaries.

Implicit boundary conditions cannot currently be used for multi-species flows, or if the IMPLICIT OVERFLOW keyword is specified.

 $See\ Also:$ IMPLICIT, DEBUG 38, DEBUG 41

IMPLICIT ORDER — Order of implicit time marching

 ${\tt IMPLICIT\ ORDER\ [\underline{ONE}\,|\,TWO]\ [\it zone_selector]}$

This keyword sets the order of the implicit time marching scheme. The default is first order. When TWO is specified, second-order time marching will be used.

Second-order time marching may be used with both structured and unstructured grids, and may be used in combination with Global Newton time marching. With structured grids, it may be used with any implicit operator except MacCormack's modified approximate factorization procedure (IMPLICIT MACCORMACK).

See Also: IMPLICIT, NEWTON, TEMPORAL

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INCLUDE — Include a file in the standard input

 ${\tt INCLUDE}\ \mathit{filename}$

This keyword allows the contents of an external file to be incorporated into Wind-US's standard input, just as if they were part of the input data (.dat) file. The *filename* must be either the full path name for the file, or a relative path name within the directory in which Wind-US is being run.

"Include" files may be nested.

INITIALIZE — Initialize in grid direction

Structured Grids

 ${\tt INITIALIZE~[ALONG]~[\underline{PLUS}|MINUS]~\{I|J|K\}~[LINES]~[} {\it zone_selector} {\tt INITIALIZE~[ALONG]~[} {\it value} {\it v$

By default, in a new (i.e., non-restart) run, the direction for the initial velocity is determined by the information specified with the FREESTREAM and ARBITRARY INFLOW keywords. For structured grids, the INITIALIZE keyword may be used to specify that the initial velocity is in the $\pm i$, $\pm j$, or $\pm k$ direction. The magnitude of the initial velocity is unchanged.

Note that one of I, J, or K must be specified, and that the PLUS direction is the default.

See Also: FREESTREAM, ARBITRARY INFLOW

ITERATIONS — Set number of iterations per cycle

Structured Grids

```
{ITERATIONS [PER] [CYCLE] | ITER_CYCLE} niters [PRINT [FREQUENCY] freq] \
   [zone_selector]
```

Unstructured Grids

```
{ITERATIONS [PER] [CYCLE] | ITER_CYCLE} niters [zone_selector]
```

This keyword allows the user to control the number of iterations performed in each zone. The parameter niters specifies the number of iterations per cycle. The default value is 5 for structured grids and 1 for unstructured grids. In addition, niters may have the following special values:

- The indicated zone(s) will be bypassed, but the zone coupling will still take place. This is not a good idea as the active zone will update inactive (n=0) zones, overwriting the existing data. This inactive zone will then pass the active zone's data back to it.
- Perform no iterations in the indicated zone and do not update this zone with adjacent zone information (but do pass this zone's frozen information to other zones each cycle).
- -2Perform no iterations in the indicated zone and do not update this zone with adjacent zone information (and do not pass this zone's frozen information to other zones).

For structured grids, residuals will be written to the .lis file every freq iterations, but only for cycles consistent with the PRINT FREQUENCY specified with the CYCLES keyword. The default value for freq is 1.

Note that the print frequency does not affect integrated properties specified using the LOADS keyword block. The print frequency for integrated properties is controlled by the PRINT keyword in the LOADS keyword block.

The zone_selector, if specified, must appear last. The print frequency, however, is global; the value specified on the last ITERATIONS entry in the input data (.dat) file will be used in all zones.

Example

For a structured grid, if the user specifies

```
CYCLES 1000 PRINT FREQUENCY 10
ITERATIONS PER CYCLE 10 PRINT FREQUENCY 5
```

cycle time information will be printed for cycles 1, 11, 21, etc. Residuals will be printed for iterations 10 (in cycle 1), 110 (cycle 11), 210 (cycle 21), etc.

See Also: CYCLES, LOADS, NAVIER-STOKES ITERATIONS, TURBULENCE

LAX DAMPING — Dissipation parameters for explicit differencing scheme

Unstructured Grids

```
LAX [DAMPING] [EXPLICIT dmp1] [IMPLICIT dmp2] [zone\_selector]
```

This keyword may be used to specify scalar dissipation parameters for the explicit differencing scheme for unstructured grids. The value dmp1 is the flux dissipation parameter, and dmp2 is the Jacobian dissipation parameter. The flux dissipation parameter only applies to the Lax (Rusanov) scheme, while the Jacobian dissipation parameter applies to all three schemes (i.e., Rusanov, Roe, and HLLE).

The default values are 0.25 and 1.5, respectively.

See Also: CONVERGE LOAD, PRECOND, RHS, DEBUG 11, DEBUG 23

LOADS — Flowfield integration (block)

Structured Grids

```
LOADS

[PRESSURE [OFFSET] {FREESTREAM | val}]

PRINT [PLANES] [TOTALS] [ZONES] [LIFT | DRAG] [FREQUENCY freq] [MAKE_FRC file]

[REFERENCE AREA aref]

[REFERENCE LENGTH lref]

[REFERENCE MOMENT [CENTER] xc yc zc]

ZONE nzn

SURFACE {I val | J val | K val} word1 [word2 [word3...]]

SUBSET I range J range K range word1 [word2 [word3...]]

ENDLOADS
```

Unstructured Grids

```
LOADS

[PRESSURE [OFFSET] {FREESTREAM | val}]

PRINT [PLANES] [TOTALS] [ZONES] [LIFT | DRAG] [FREQUENCY freq] [MAKE_FRC file]

[REFERENCE AREA aref]

[REFERENCE LENGTH lref]

[REFERENCE MOMENT [CENTER] xc yc zc]

ZONE nzn

SURFACE {I|U} val word1 [word2 [word3...]]

ENDLOADS
```

Flowfield properties may be integrated during the course of a Wind-US run to check convergence and solution quality. The values printed to the list file represent the force coefficients F/A_rq_r and moment coefficients $M/L_rA_rq_r$ in the x, y, and z coordinate directions, where F and M are the force and moment, and L_r , A_r , and q_r are the reference values of length, area, and dynamic pressure. Integrated mass and momentum fluxes may also be computed. The values are written into the list output (.lis) file.

The various elements of the LOADS keyword block are defined below. Unless noted otherwise, the keywords apply to both structured and unstructured grids.

```
PRESSURE [OFFSET] {FREESTREAM | val}
```

This keyword controls the pressure integration. By default, the code uses $P-P_{\infty}$ in all pressure integrations, where P_{∞} is the static pressure at freestream conditions. This keyword permits the user to specify the pressure offset value, such that:

```
FREESTREAM Use P-P_{\infty} val \qquad \qquad \text{Use } P-val \text{PRINT [PLANES] [TOTALS] [ZONES] [LIFT | DRAG] [FREQUENCY freq] [MAKE_FRC file]}
```

This keyword controls the output data from the integration. Options are turned 'on' by including the appropriate keyword.

PLANES Output the result of the integration for each surface and/or subset specified.

TOTALS Output integration grand totals over all the zones at the end of each cycle.

ZONES Output integration totals for each zone.

LIFT | DRAG Output lift, drag, and side forces instead of (x, y, z) force components. Directions

for the lift, drag, and side force components are computed from the angles of

attack and yaw specified using the FREESTREAM keyword.

FREQUENCY Output integration results (except for grand totals) every freq iterations. The

default value is 5.

MAKE_FRC Write grand totals to the summary file file at the end of each cycle, in addition

to the .lis file. This requires that PRINT TOTALS also be specified.

REFERENCE AREA aref REFERENCE LENGTH lref

REFERENCE MOMENT [CENTER] $xc\ yc\ zc$

These keywords specify the reference area aref in square inches; the reference length lref in inches; and the coordinates (xc,yc,zc) of the reference moment center in inches. The default values are 1.0 for aref, 1.0 for lref, and (0.0, 0.0, 0.0) for (xc, yc, zc).

ZONE nzn

This keyword specifies the zone number nzn for the surfaces and/or subsets which follow.

SURFACE {I val | J val | K val} word1 [word2 [word3...]]

This form of the SURFACE keyword applies to structured grids, and specifies the coordinate surface in zone nzn over which the integration is to be performed. The parameter val is a coordinate index specifying the surface.

The *word* parameters are keywords specifying what flowfield properties are to be computed, plus a couple of additional control values. The following keywords are currently available.

FORCE Compute pressure forces.

MOMENT Compute moments.

VISCOUS Also compute viscous forces (when FORCE is specified) and moments (when

MOMENT is specified).

MASS Compute mass flow.

MOMENTUM Compute momentum.

NORMAL inorm inorm = ± 1 , specifying the desired normal vector direction; +1 for the increas-

ing index direction and -1 for the decreasing index direction. *inorm* defaults toward the interior of the grid if the integration surface is on a boundary of the

zone.

NOSLIP Integrate only over points where the total velocity is 0. [Currently (Wind-US

3.88), the NOSLIP option only works when specified with SUBSET, not SURFACE.

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At least one flowfield property must be specified. For integration surfaces that are not part of a zonal boundary, the normal direction must also be specified.

Note that when VISCOUS is specified, the viscous forces and moments are printed to the .lis file as separate values; they are not summed with the pressure forces and moments. Note also that you can't compute just the viscous values. E.g., to get the viscous forces you must also get the pressure forces. Specifying just SURFACE J 1 VISCOUS will result in no forces being printed to the .lis file for the j=1 surface.

```
SURFACE {I|U} val word1 [word2 [word3...]]
```

This form of the SURFACE keyword applies to unstructured grids, and specifies the unstructured surface in zone nzn over which the integration is to be performed. The parameter val is the surface number.

The word parameters are keywords specifying what flow field properties are to be computed. The following keywords are currently available.

FORCE Compute pressure forces.

MOMENT Compute moments.

VISCOUS Also compute viscous forces (when FORCE is specified) and moments (when MOMENT

is specified).

MASS Compute mass flow.

MOMENTUM Compute momentum.

HEAT Compute heat flux. This keyword is only available for isothermal walls.

At least one flowfield property must be specified.

```
SUBSET I range J range K range word1 [word2 [word3...]]
```

This keyword may be used with structured grids to define a subset of a computational surface in zone nzn over which the integration is to be performed. The range parameters take one of the following forms:

index1 index2 Starting and ending indices in the specified direction. LAST may be used for the last index.

ALL Equivalent to 1 LAST.

For three-dimensional cases, the starting and ending indices for one (and only one) of the I, J, or K parameters must be the same. For two-dimensional cases, the K parameter must be specified as either K 1 1 or K ALL; one or both of the I and J parameters may have different starting and ending indices.

The word parameters are the same as those described above for the SURFACE keyword for structured grids.

Example

```
LOADS
```

PRESSURE OFFSET 0.0

PRINT PLANES ZONES TOTALS LIFT FREQUENCY 2

```
REFERENCE AREA 100.

REFERENCE LENGTH 12.

REFERENCE MOMENT CENTER 35. 36. 78.

ZONE 4

SUBSET I ALL J 1 1 K 15 LAST FORCE MOMENT VISCOUS NOSLIP

ZONE 10

SURFACE I 1 MOMENTUM MASS

ENDLOADS
```

 $See \ Also:$ CONVERGE LOAD, FIXED_CL, POSTPROC, WALL FUNCTION

MARCHING — Parabolized Navier-Stokes algorithm

Structured Grids

MARCHING [LIMITER value] [CHECKPOINT interval] [COPY]

For structured grids, this keyword enables Wind-US's spatial marching, or parabolized Navier-Stokes (PNS), algorithm for flowfields which are supersonic in the computational i-direction. In this mode, Wind-US marches from the i=1 to the $i=i_{max}$ computational surface, attempting to compute a steady-state solution at each plane before moving on to the next one. Using the PNS algorithm significantly reduces the computing time required for supersonic solutions.

This keyword enables the marching limiter, which limits the change in LIMITER value the solution vector \mathbf{Q} to $(value)\mathbf{Q}$. I.e.,

 $\Delta \mathbf{Q} < (value)\mathbf{Q}$

CHECKPOINT interval The parameter *interval* specifies the number of *i*-planes to be computed

before writing the current flowfield to the solution file. The default is 10.

COPY This keyword requests that Wind-US copy the solution from the most

recently computed i-plane to the upcoming i-plane, giving a (hopefully) better initialization to the new i-plane than simply starting from

freestream flow.

Notes

• Marching is only available for structured grids.

- Marching is only available when Wind-US is run in serial mode.
- Marching is not available with the one-equation turbulence models (BARTH, SPALART).
- Marching requires that one of the following explicit operators be used. (See the RHS keyword.)
 - Coakley (any order)
 - Roe (first-order upwind, second-order upwind, third-order upwind-biased, first-order upwind modified for stretched grids, or second-order upwind-biased modified for stretched grids)
 - Van Leer (first-order upwind, second-order upwind, or third-order upwind-biased)
- The Roe second-order upwind-biased explicit operator modified for stretched grids uses the i+1 grid point, which is invalid in a PNS solution. When this operator is used, Wind-US automatically changes to first order in the *i*-direction.

See Also: RHS, TURBULENCE

MASS FLOW — Outflow boundaries, mass flow

Structured Grids

```
MASS [FLOW] {RATE [ACTUAL | CORRECTED] | RATIO} value \setminus [PRESSURE | DIRECT [RELAXER <math>rlxr]] [ORDER {ZERO|0|ONE|1}] \ [ZONE] range1[, range2[, ... rangen]]
```

Unstructured Grids

```
MASS [FLOW] {RATE [ACTUAL | CORRECTED] | RATIO} value \setminus [ZONE] \ range1 [, range2 [, ... rangen]]
```

This keyword allows the user to specify mass flow at outflow boundaries in the flowfield.

Structured Grids

The input keywords applicable to structured grids are as follows:

RATE value represen

 $\it value$ represents the mass flow rate in $\it lb_m/sec$, and may be actual (the default) or corrected, as specified by the ACTUAL or CORRECTED keyword. The specified $\it value$ must be positive. The corrected air flow is defined as

$$W_c = W_{actual} \frac{\theta_x^{0.5}}{\delta_x}$$

where

$$\delta_x = P_x/P_0 \qquad \theta_x = T_x/T_0$$

and P_x and T_x are the total pressure and temperature at the duct exit, and P_0 and T_0 are equal to 14.7 psi and 520 °R, respectively.

RATIO value represents the mass flow ratio. Again, the specified value must be positive. The actual mass flow is computed as

$$\dot{m} = (value)\rho_{\infty}U_{\infty}A_{cap}$$

where A_{cap} is the capture area found in the .cgd file zonal parameters.

PRESSURE A spatially-constant pressure is set at the boundary, and modified as the solution proceeds until the desired mass flow is achieved. This is the default.

DIRECT The momentum, and thus the mass flow, is modified directly, and the pressure adjusts as the solution proceeds.

RELAXER The specified mass-flow rate will be relaxed using the relaxation factor rlxr. This option only applies when DIRECT is specified. The default value for rlxr is 1.0 (i.e., no relaxation).

ORDER Either zeroth- or first-order extrapolation will be used, as specified. The default is zeroth-order.

In the zone specification, the range parameter(s) must be one of the following forms:

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zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

Note that if multiple zones are listed, the specified mass flow will be applied in each zone.

With the PRESSURE option, the pressure will be constant over the entire outflow boundary, resulting in poor solutions for flows that should have cross-flow pressure gradients in that region. With the DIRECT option, cross-flow pressure gradients may be present at the outflow boundary, and the mass flow will be equal to the user-specified value (for rlxr = 1) for all iterations.

For flows with negligible cross-flow pressure gradients, the results and convergence rates using the PRESSURE and DIRECT options are nearly the same. For a test case with a significant cross-flow pressure gradient near the outflow boundary, the computed pressures using the two options differed by as much as 10%. The PRESSURE option, although non-physical for this case, had a slightly better convergence rate.

Internally, to apply this boundary condition Wind-US does the following:

- 1. At all the boundary points, the density, momentum, and energy are extrapolated from the interior to the boundary using either zeroth- or first-order extrapolation, as specified.
- 2. The mass flow at the boundary is computed by numerical integration.
- 3. If PRESSURE was specified, a new downstream pressure is computed based on the difference between the computed and specified mass flow rates.
- 4. If DIRECT was specified, a mass flow correction factor is computed as

$$f_{corr} = 1 + r \left(\frac{\dot{m}_{spec}}{\dot{m}_{int}} - 1 \right)$$

where r is the input relaxation factor rlxr, and \dot{m}_{spec} and \dot{m}_{int} are the user-specified and integrated mass flow rates, respectively.

- 5. For each boundary point, the density and momentum, plus the pressure, effective gamma, compressibility factor, and speed of sound, are extrapolated from the interior to the boundary using either zeroth- or first-order extrapolation, as specified.
- 6. If DIRECT was specified, the extrapolated momentum values at the outflow boundary are modified using

$$(\rho u) = f_{corr}(\rho u)_{ext}$$

 $(\rho v) = f_{corr}(\rho v)_{ext}$

$$(\rho w) = f_{corr}(\rho w)_{ext}$$

- 7. If PRESSURE was specified, the energy at each boundary point is computed, consistent with the downstream pressure value from step 3 and the extrapolated values of density, etc., from step 5.
- 8. If DIRECT was specified, the energy at each boundary point is computed, consistent with the momentum values from step 6 and the extrapolated values of density, pressure, etc., from step 5.

Extrapolation Notes

The default for all extrapolation is zeroth-order (i.e., conditions at the boundary are set to the values at the computational plane adjacent to the boundary). This results in a discontinuous slope in

flow values near the outflow boundary, which may be important for flows with significant streamwise pressure gradients. First-order extrapolation yields smoother results.

For flows with little or no streamwise pressure gradient near the outflow boundary, the results using zeroth- and first-order extrapolation are essentially identical. Convergence rates and the final residual values are generally better with zeroth-order extrapolation, however, so the default zeroth-order extrapolation is recommended.

For flows with significant streamwise pressure gradients near outflow boundaries, zeroth-order extrapolation can give poor results at the outflow boundary, and in some cases these can affect values at the inflow boundary. First-order extrapolation is thus recommended for these flows.

Examples

```
MASS FLOW RATIO 0.95 ZONE 2
MASS FLOW RATE ACTUAL 180. ZONE 3
MASS FLOW RATE CORRECTED 220. ZONE 4
```

See Also: COMPRESSOR FACE, DOWNSTREAM PRESSURE, DOWNSTREAM MACH, TEST 123, TEST 153, TEST 160

Unstructured Grids

The input keywords applicable to unstructured grids are as follows:

RATE

 $\it value$ represents the mass flow rate in $\it lb_m/sec$, and may be actual (the default) or corrected, as specified by the ACTUAL or CORRECTED keyword. The specified $\it value$ must be positive. The corrected air flow is defined as

$$W_c = W_{actual} \frac{\theta_x^{0.5}}{\delta_x}$$

where

$$\delta_x = P_x/P_0$$
 $\theta_x = T_x/T_0$

and P_x and T_x are the total pressure and temperature at the duct exit, and P_0 and T_0 are equal to 14.7 psi and 520 °R, respectively.

RATIO

value represents the mass flow ratio. Again, the specified value must be positive. The actual mass flow is computed as

$$\dot{m} = (value)\rho_{\infty}U_{\infty}A_{cap}$$

where A_{cap} is the capture area found in the .cgd file zonal parameters.

In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

Note that if multiple zones are listed, the specified mass flow will be applied in *each* zone.

For unstructured grids, a spatially-constant pressure is set at the boundary, and modified as the solution proceeds until the desired mass flow is achieved. No DIRECT option is available. In addition, the extrapolation for unstructured grids is zeroth-order.

See Also: DOWNSTREAM PRESSURE, TEST 123, TEST 153, TEST 160

MFD — Magneto-Fluid Dynamics Model (block)

Structured Grids

```
[OUTPUT {BFIELD | CONDUCTIVITY | CURRENT | EFIELD | VOLTAGE | LORENTZ}]
   [RELAX_MFD nriter]
   [UPDATE nuiter]
   [RADIATION emiss lref [tback]]
   {LORENTZ {CFL | FORCE dummy FREQUENCY f [DUTY du] [SCALE sc] \
                    PHASES n PATTERNS p1 p2 p3 ... pn}
    CONDUCTIVITY {CFL | \
                   CONSTANT sigma | \
                   EQUILIBRIUM {ARGON | AIR | GAS} [POTASSIUM mk] | \
                   LINEAR t1 \ sig1 \ t2 \ sig2 \ | \ \setminus
                   PREDICTED [USING] [LIN-RESSLER | BOEING]}
    BFIELD {CFL | CONSTANT bz | BLOCKS nbblocks}
    {EFIELD {CFL | CONSTANT ey | BLOCKS neblocks} | \
     VOLTAGE {CFL | BOUNDARIES nvbnd | PARAMETERS mitvlt vlttol vltrx vltry vltrz vltfac}}
    EXTERNAL [INPUT] [MODE] PEM}
ENDMFD
```

The MFD keyword block allows the user to include body force source terms in the Navier-Stokes equations resulting from magneto-fluid dynamics effects. This capability is only available for structured grids.

Control Functions

MFD

Defines the beginning of the MFD block.

ENDMFD

Defines the end of the MFD block.

```
OUTPUT {BFIELD | CONDUCTIVITY | CURRENT | EFIELD | VOLTAGE | LORENTZ}
```

The specified data (i.e., magnetic field, conductivity, current density, electric field, voltage, or Lorentz force) will be written into into the flow (.cfl) file. Multiple OUTPUT keywords may be specified, to write multiple types of data into the .cfl file. When VOLTAGE BOUNDARIES is specified (see below), the current density, electric field, and voltage are automatically written into the .cfl file.

```
{\tt RELAX\_MFD} \ \ nriter
```

The MFD source terms in the Navier-Stokes equations will be relaxed over *nriter* iterations. The default for *nriter* is 1.

```
{\tt UPDATE} \ nuiter
```

The MFD source terms in the Navier-Stokes equations will be updated every *nuiter* Navier-Stokes iterations. The default for *nuiter* is 1.

If VOLTAGE BOUNDARIES is used, *nuiter* must be 1 (the default). In this case, consider running a few iterations with VOLTAGE BOUNDARIES to compute the electric field, then restarting using EFIELD CFL instead of VOLTAGE BOUNDARIES.

```
RADIATION emiss\ lref\ [tback]
```

Estimate the energy loss due to thermal radiation of the fluid with emissivity *emiss*, optical depth lref, and background temperature tback (°R). The default for tback is the freestream static temperature.

Body Force Determination

The body force resulting from the MFD terms can be added in one of three ways: (1) by directly specifying the Lorentz force; (2) by specifying the conductivity, the magnetic field, and either the electric field or voltage; or (3) by reading the data from the .cfl file, stored there using an external solver.

Specifying the Lorentz Force

The following keyword is used to directly specify the Lorentz force. If this method is used, the CONDUCTIVITY, BFIELD, EFIELD, VOLTAGE, and EXTERNAL keywords are not allowed.

```
LORENTZ {CFL | FORCE dummy FREQUENCY f [DUTY du] [SCALE sc] \ ----PHASES n PATTERNS p_1, p_2, ..., p_n}
```

CFL Read the Lorentz force field directly from the flow (.cfl) file FORCE dummy N_L Lorentz force distributions are stored in the flow (.cfl) file, defining a time-dependent cyclical force field. For each distribution the variable names for the Lorentz force components are Lxi, Lyi, and Lzi, where i varies from 1 to N_L . The appropriate distribution is read and used, based on the current integrated time and the specified phasing information, then scaled and applied. [Currently, due to a coding problem, a value dummy must be specified after the word FORCE, but is not otherwise used.] FREQUENCY fNumber of cycles/second for the Lorentz force field DUTY duFraction of each phase in which the Lorentz force will be applied. The default value is 1.0. SCALE sc Scale factor. The Lorentz force will be multipled by sc before adding it to the equations. The default value is 1.0. PHASES nNumber of pattern phases in a cycle PATTERNS p_1 , p_2 , ..., p_n The Lorentz force distribution to use in each phase. A total of n values must be specified, where n is the number of pattern phases specified with PHASES. For each phase, the value p_i is an integer,

.cfl file to be applied for that phase.

from 1 to N_L , defining the particular force distribution from the

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Specifying the MFD Fields

If the MFD fields are being specified, you must use the CONDUCTIVITY and BFIELD keywords to specify the conductivity and the magnetic field, and either the EFIELD or VOLTAGE keywords to specify the electric field or voltage.

```
CONDUCTIVITY {CFL | \ CONSTANT sigma | \ EQUILIBRIUM {ARGON | AIR | GAS} [POTASSIUM mk] | \ LINEAR t1 sig1 t2 sig2 | \ PREDICTED [USING] [LIN-RESSLER | BOEING]}
```

This keyword specifies the electrical conductivity in mhos/meter.³⁵

CFL Read the conductivity from the flow (.cfl) file

CONSTANT sigma Hold the conductivity constant at the value sigma

EQUILIBRIUM {ARGON | AIR | GAS} [POTASSIUM mk]

Estimate the electron density as a function of temperature for the indicated gas as input to the Lin & Ressler conductivity model. The default gas is air. If POTASSIUM mk is specified, the effect of the mass

fraction mk of potassium will be included.

LINEAR t1 sig1 t2 sig2 Set the conductivity to sig1 at and below the temperature t1; to sig2

at and above the temperature t2; and use a linear distribution for

temperatures between t1 and t2. The temperatures are in ${}^{\circ}R$.

PREDICTED [USING] [LIN-RESSLER | BOEING]

Use real-gas predicted electron densities for input to the indicated conductivity model. The default is the Lin & Ressler model.

BFIELD {CFL | CONSTANT bz | BLOCKS nbblocks}

This keyword specifies the magnetic field in tesla.

CFL Read the electric field from the flow (.cfl) file

CONSTANT bz Hold the magnetic field constant at the value bz, in the z coordinate direction

BLOCKS nbblocks Specify the magnetic field by reading in nbblocks blocks of data containing the

magnetic field vector at selected coordinate points. The data immediately follows the $\tt BFIELD\ BLOCKS\ keyword$. See the $\tt Field\ Block\ Description$ for

details and an example.

EFIELD {CFL | CONSTANT ey | BLOCKS neblocks}

This keyword specifies the electric field in Volts/meter.

CFL Read the electric field from the flow (.cfl) file

CONSTANT ey Hold the electric field constant at the value ey, in the y coordinate direction

 $^{^{35}\}mathrm{The}$ keyword SIGMA may be used as an alias for CONDUCTIVITY.

BLOCKS neblocks

Specify the electric field by reading in *neblocks* blocks of data containing the electric field vector at selected coordinate points. The data immediately follows the EFIELD BLOCKS keyword. See the **Field Block Description** for details and an example.

With this keyword the electric field is determined by specifying the electric potential.

CFI.

Read the electric potential field from the flow (.cfl) file

BOUNDARIES nvbnd

Specify the electric potential at *nvbnd* zonal boundary regions. The data immediately follows the VOLTAGE BOUNDARIES keyword. See the Voltage Boundary Description for details and an example.

PARAMETERS mitvlt vlttol vltrx vltry vltrz vltfac

Iterate a maximum of *mitvlt* iterations (the default is 10,000) to a tolerance of *vlttol* (a positive value means to a level of *vlttol*, a negative value means |vlttol| orders of magnitude; the default is a level of 10^{-12}) with implicit factors *vltrx*, *vltry*, and *vltrz* (the default values are 1.0, currently these must all be set to 0.0) and with an over-relaxation factor of *vltfac* (the default is 1.0, but a more typical value is 0.4).

Note that when VOLTAGE BOUNDARIES is specified, you must also use (separately) VOLTAGE PARAMETERS to specify the iteration controls for solution of the electric potential equation, even though they have default values. That's because the defaults for *vltrx*, *vltry*, and *vltrz* are all 1.0, but only 0.0 is currently allowed.

Using an External Solver

```
EXTERNAL [INPUT] [MODE] PEM
```

This keyword indicates that the MFD fields have been computed using external solver, and should be read from the .cfl file.

Field Block Description

By using the BFIELD BLOCKS and/or EFIELD BLOCKS keywords, the magnetic and/or electric field may be determined from blocks containing the field vector at selected coordinate points. Each block consists of eight points in space, with the corresponding field vector at each of those points. The region contained within the blocks is filled using tri-linear interpolation between the specified points. Up to eight blocks may be specified for each field type. The spatial locations of the blocks may overlap, with the later-specified blocks overwriting the earlier ones.

There are eight lines of input per block, one for each of the eight coordinate points. Each line contains six values—the x, y, and z coordinates, and the field vector components in the x, y, and z directions.

The following example specifies the magnetic field using two blocks. (The comments in a slanted font are not part of the input.)

					OCKS 2	BFIELD BLO
Upstream plane of block 1	0.0	0.0	0.0	0.0	0.0	60.0
	0.0	0.0	0.0	0.0	20.0	60.0
	0.0	0.0	0.0	1.0	0.0	60.0

	0.0	0.0	0.0	1.0	20.0	60.0
Downstream plane of block 1	10.0	0.0	0.0	0.0	0.0	120.0
	5.0	0.0	0.0	0.0	20.0	120.0
	10.0	0.0	0.0	1.0	0.0	120.0
	5.0	0.0	0.0	1.0	20.0	120.0
$Upstream\ plane\ of\ block\ 2$	10.0	0.0	0.0	0.0	0.0	120.0
(same as downstream of 1)	5.0	0.0	0.0	0.0	20.0	120.0
	10.0	0.0	0.0	1.0	0.0	120.0
	5.0	0.0	0.0	1.0	20.0	120.0
Downstream plane of block 2	0.0	0.0	0.0	0.0	0.0	180.0
	0.0	0.0	0.0	0.0	20.0	180.0
	0.0	0.0	0.0	1.0	0.0	180.0
	0.0	0.0	0.0	1.0	20.0	180.0
	B_z	B_y	B_x	\boldsymbol{z}	y	\boldsymbol{x}

Voltage Boundary Description

Voltage boundaries are used to specify the boundary conditions at zone boundaries for the electric potential solver. The electric potential solver will take into account gradients and discontinuities in conductivity, as well as the electromotive force (EMF) induced by the fluid motion through the magnetic field. Currently, accurate voltage predictions require a grid with low skewness.

Up to 64 boundaries may be specified. The conditions at each boundary are specified on a single line, with eight values—zone, face, type, L1, L2, M1, M2, value—defined as follows:

zone	Zone containing the boundary
face	The boundary face, specified as a number from 1 to 6 corresponding to the $i_1, i_{max}, j_1, j_{max}, k_1$, and k_{max} face, respectively
type	The boundary type, specified as 0 for a reflection boundary, and 1 for specified voltage
L1, L2, M1, M2	The indices on the face over which the boundary condition applies, as follows

Indices
$j_{low}, j_{high}, k_{low}, k_{high}$
$k_{low}, k_{high}, i_{low}, i_{high}$
$i_{low},i_{high},j_{low},j_{high}$

valueThe voltage for specified-voltage boundaries; 0.0 for reflection boundaries

The following example specifies eight voltage boundaries. (The comments in a slanted font are not part of the input.)

VOLTAGE	BOUN	DARIES	8				
1	1	0	1	30	1	1	0.0
1	1	1	31	61	1	1	0.0
1	2	1	1	31	1	1	10000.0
1	2	0	32	61	1	1	0.0
1	3	0	1	1	1	30	0.0
1	3	1	1	1	31	61	10000.0
1	4	1	1	1	1	31	0.0
1	4	0	1	1	32	61	0.0

Zone Face Type L1 L2 M1 M2 Value

MOVING WALL — Specify moving wall boundaries

```
MOVING [WALL] region \setminus \{TRANSLATE \{I \ speed \mid \{X|Y|Z\} \ speed \mid u \ v \ w\} \mid \setminus SPIN \{\{X|Y|Z\} \ \omega \ c_1 \ c_2 \mid x_c \ y_c \ z_z \ \{ANGLES \ \theta \ \phi \ \psi \mid AXIS \ \omega \ x_a \ y_a \ z_a\}\} \mid \setminus TANGENT_ONLY\}
```

This keyword may be used to define a translating and/or spinning no-slip wall. Moving walls are activated in regions specified as bleed in the .cgd file (set with GMAN), where region is the "bleed" region number.

There are three options for specifying a translating wall.

TRANSLATE I Wall is moving in the i-direction at the velocity speed, in feet per second.

This is only available for structured grids.

TRANSLATE $\{X|Y|Z\}$ Wall is moving in physical space in the x, y, or z direction at the

velocity *speed*, in feet per second.

TRANSLATE $u \ v \ w$ Wall is moving in physical space, where u, v, and w are the velocities

in the x, y, and z directions, in feet per second.

There are two options for specifying a spinning wall.

SPIN $\{X|Y|Z\}$ ω c_1 c_2 Wall is spinning in physical space about the x, y, or z axis, with rotation

rate ω in degrees per second. For rotation about the x axis, c_1 and c_2 are the coordinates of the center of rotation in the y-z plane. Similarly, for rotation about the y axis, they're the coordinates of the center of rotation in the x-z plane, and for rotation about the z axis, they're the

coordinates of the center of rotation in the x-y plane.

SPIN x_c y_c z_c {...} Wall is spinning in physical space, with the coordinates x_c , y_c , and z_c

defining the center of rotation. The rate and direction of rotation may be specified using either the ANGLES or AXIS keyword. For ANGLES, θ , ϕ , and ψ specify the rotation rates about the x, y, and z axes, in degrees per second. For AXIS, ω is the rotation rate in degrees per second, and x_a , y_a , and z_a are the x, y, and z coordinates that, along with x_c , y_c ,

and z_c , define the axis of rotation.

Separate MOVING WALL keywords may be used to specify a single *region* that is both translating and spinning in physical space. The TRANSLATE I option, however, may not be specified for a *region* that is also moving in physical space.

An additional MOVING WALL keyword specifying TANGENT_ONLY may be used to force the flow to be tangential to the wall by removing the normal velocity components. Note that this only makes sense if a separate MOVING WALL keyword (or keywords) is used to specify a translating and/or spinning wall.

${\bf NAVIER\text{-}STOKES\ ITERATIONS-Navier\text{-}Stokes\ sub-iterations}$

NAVIER-STOKES ITERATIONS iter [zone_selector]

Wind-US organizes the equations to be solved into logical "groups" that are solved together. It also allows multiple iterations of a specific group (i.e., sub-iterations) for each "iteration per cycle". (The number of iterations per cycle is set using the ITERATIONS keyword.)

The NAVIER-STOKES ITERATIONS keyword allows the user to specify the number of sub-iterations for the Navier-Stokes equation group (which includes any chemical species equations) performed in each zone for each "iteration per cycle". The default value is one, indicating that each "iteration per cycle" corresponds to one iteration of the Navier-Stokes equations. Note that if *iter* is set to zero, the Navier-Stokes equations will not be solved at all.

See Also: ITERATIONS, TURBULENCE

NEWTON — Use Global Newton time stepping

NEWTON [TIME LEVELS ntlvls] [CONVERGE {LEVEL | $\underline{\texttt{ORDER}}$ } newcvg]

This keyword may be used to specify that the Global Newton time stepping algorithm is to be used. The parameter *ntlvls* specifies the number of Global Newton time levels to advance. The default value is 30.

There are two possible procedures for determining the overall global convergence.

CONVERGE LEVEL Convergence is assumed when

$$|\mathbf{Q}^n - \mathbf{Q}^{n-1}| < newcvg$$

where \mathbf{Q} represents the vector of dependent variables, and n is the Newton time level.

CONVERGE ORDER Convergence is assumed when

$$\frac{|\mathbf{Q}^n - \mathbf{Q}^{n-1}|}{\max(|\mathbf{Q}^n - \mathbf{Q}^{n-1}|)} < 10^{-newcvg}$$

The default is CONVERGE ORDER 3.

Within a Newton time level, the number of cycles and iterations to be run is specified using the CYCLES and ITERATIONS keywords.

The convergence criteria within a Newton time level may be specified using the CONVERGE keyword. Note that for structured grids the default for the CONVERGE keyword is a four-order of magnitude decrease in the maximum residual. If Newton iteration is being used for a steady flow problem with structured grids, with the default of three orders of magnitude for the global convergence criteria, it would make sense to also use three orders of magnitude for the convergence criteria within a Newton time step.

See Also: CFL, CYCLES, ITERATIONS, CONVERGE, IMPLICIT ORDER, TEMPORAL

${\bf OUTFLOW\ NON-REFLECTING-Outflow\ boundaries,\ non-reflecting}$

Structured Grids

```
OUTFLOW [NON-REFLECTING] [ZONE] range1[,range2[, ... rangen]]
```

This keyword imposes a non-reflecting, subsonic outflow boundary condition at outflow boundaries in structured zones. Acoustic disturbances reaching the boundary are essentially eliminated. It is actually implemented using the Paynter compressor face model (Slater and Paynter, 2000), but with the response coefficients α and β set to zero.

In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

This boundary condition is intended to be used for time-accurate flows with a CFL number less than one; however, it may be applicable in certain cases for steady-state simulations.

See Also: COMPRESSOR FACE PAYNTER, TEST 123

PERIODIC — Periodic boundaries

Periodic boundaries are currently supported only for structured grids. They are treated as normal coupled boundaries, with the connection data stored in the grid (.cqd) file. The CONNECT MODE option in GMAN is used to specify how to move one boundary surface to overlay it on the corresponding periodic boundary surface. There are two connection modes — translation and rotation. For translation, the user defines the Δx , Δy , and Δz movement for the surface. For rotation the user sets the center of rotation and the angles. GMAN then moves the current boundary surface using the specified method, and attempts to couple the two surfaces together as if they were coexistent boundaries. If successful, it saves the data as normal coupling data for Wind-US. Thus the boundaries look connected even though they are physically separated. The periodic boundary surfaces do not have to be in the same zone, nor do they have to be point matched. They only have to line up physically once the movement has been performed.

Specifically, to set a periodic boundary condition for a structured grid in GMAN:

- In graphics mode, from the main menu select BOUNDARY COND.
- Pick the zone and boundary for one of the two periodic boundaries.
- If the boundary condition at the boundary is not UNDEFINED (you may select IDENTIFY POINTS to check), change it to UNDEFINED by doing:
 - Select MODIFY BNDY.
 - Select CHANGE ALL.
 - Select UNDEFINED.
 - From the main menu, select BOUNDARY COND.
 - Select YES-UPDATE FILE.
- Select MODIFY BNDY.
- Select COUPLE.
- Select SEL OTHER BND.
- Pick the zone and boundary for the other periodic boundary. (Note that no default zone is pre-selected here, even for single-zone grids, so you must select both the zone and boundary.)
- Select SET COUP MODE.
- Under CONNECT MODE, click on the "** NONE **".
- Respond to the prompts at the bottom of the screen to set the connection mode, and to specify the translation or rotation data.
- Select CONNECT (not COUPLE) to actually connect the two boundaries.

Another procedure is available for setting up a periodic boundary condition within a single structured zone, and may be useful for cases with higher order differencing schemes. See the description of TEST 72 and TEST 73 for details.

POSTPROC — Create GENPLOT files for assessing convergence (block)

```
/!POSTPROC [gen_name]

/!{SETDEF | SET} set_options

/![{ZONE | ZONES} zone_list]

/![{PLANE | PLANES} plane_list]

/![{DIRECTION | DIR} direction]

/![{COMPONENT | COMP} [NOPRESSURE | NOVISCOUS]]

/!ENDPOSTPROC
```

The POSTPROC keyword block is used in conjunction with the <code>-genpost</code> wind script option and the LOADS keyword to automatically create GENPLOT files containing convergence information during a Wind-US run. After the Wind-US run ends, these files could then be post-processed using CFPOST to graphically assess the progress of the solution toward convergence. The <code>-genpost</code> option specifies how often the files should be updated, and the information in the <code>POSTPROC</code> block specifies what should be written into the files.

Note that every keyword in the POSTPROC block, including the opening and closing keywords, starts with the two characters "/!". If the -genpost option is not used, this keyword block is ignored.

The various elements of the POSTPROC keyword block are defined as follows:

```
/!POSTPROC [gen_name]
...
/!ENDPOSTPROC
```

The POSTPROC and ENDPOSTPROC keywords define the beginning and end of the POSTPROC block. Multiple POSTPROC blocks are allowed. The optional gen_name is the name to be used for the GENPLOT file. An extension of .gen is automatically added. The default is Postgen1 for the first POSTPROC block in the .dat file, Postgen2 for the second, etc.

```
/!{SETDEF | SET} set_options
```

The SETDEF keyword is required, and defines the quantities to be written into the GENPLOT file. These may be residuals, and/or integrated flowfield values specified using the LOADS keyword block. The values are extracted from the .lis file and written into the GENPLOT file using the utility resplt.pl in batch mode. The input string set_options contains command-line options for resplt.pl that specify the quantities to be extracted. Thus, see the "Command-Line Options" section of the resplt.pl documentation for the syntax to be used in set_options.

Note that $set_options$ is only used to define the quantities to be extracted (i.e., residuals, loads, etc.) Other options used when resplt.pl is run as a stand-alone utility, such as <code>-list</code>, <code>-plane</code>, <code>-zone</code>, etc., are automatically created by the wind scripts, and should not be specified using the SETDEF keyword.

```
/!{ZONE | ZONES} zone_list
```

Extract quantities only for the listed zones. Multiple zones and/or a range of zones may be specified using syntax like "2,3,6-8". The default is all zones.

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/!{PLANE | PLANES} $plane_list$

Extract integrated flowfield quantities only for the listed "planes". The "planes" are actually the surfaces and/or subsets specified using the SURFACE and SUBSET keywords in the LOADS keyword block. Multiple planes and/or a range of planes may be specified using syntax like "2,3,6-8". The default is all planes.

Planes are numbered in the order specified in the Wind-US input data (.dat) file, and are "global", not "zonal". Thus, the ZONE keyword in the POSTPROC block does not apply. For example, imagine a five-zone grid with two SURFACE keywords specified in the LOADS block for each zone, in order. If in the POSTPROC block you specify

ZONE 3 PLANES 1,2

you'll be extracting integrated values for the two surfaces in zone 1, not zone 3, since planes 1 and 2 are in zone 1. To extract values for the two surfaces in zone 3, you'd specify

PLANES 5,6

/!{DIRECTION | DIR} direction

When forces, moments, or momentum fluxes are being extracted, the DIRECTION keyword may be used to specify the desired direction. For details see the description of the -direction command-line option for *resplt.pl*. The default is all applicable directions.

/!{COMPONENT | COMP} [NOVISCOUS | NOPRESSURE]

When forces or moments are being extracted, only extract pressure values (when NOVISCOUS is specified) or viscous values (when NOPRESSURE is specified). The default is to extract both pressure and viscous values.

Note that viscous values are only available when VISCOUS is specified in the surface/subset specification in the LOADS keyword block. If it's not, specifying COMPONENT NOVISCOUS in the POSTPROC block will have no effect.

PRANDTL — Set the laminar and turbulent Prandtl numbers

PRANDTL Prl Prt

This keyword may be used to set the laminar and turbulent Prandtl numbers. The default values are 0.72 and 0.9, respectively.

Note that the Prandtl numbers may also be set using the GAS keyword.

 $See\ Also:\ {\tt GAS}$

PRECOND — Pre-conditioning for low Mach number flows

Unstructured Grids

PRECOND HYPERCOMP eps_{pre}

This keyword may be used to apply matrix pre-conditioning to improve (or allow) convergence for flows at low Mach numbers. While it theoretically may be used at all Mach numbers, it increases the CPU time required for each time step, and is therefore recommended for use only when the Mach number is less than about 0.05.

The input parameter eps_{pre} is an artificial compressibility factor. The recommended value is

$$eps_{pre} = \max(\sqrt{3}M_{\infty}, 0.2)$$

where M_{∞} is the freestream Mach number. Very small values for eps_{pre} will lead to very large values for artificial compressibility, and convergence problems, especially when stagnation regions are present. Values of eps_{pre} that are too large will decrease the amount of pre-conditioning, leading to slow (or no) convergence.

Additional details on the implementation of the pre-conditioning scheme, and results for various test cases, are presented by Liu and Ramakrishnan (2004).

This keyword may only be used when a physical time is specified with the CFL# keyword, not a CFL number. It also may not be used with the HLLE scheme (RHS HLLE).

Q LIMIT — Limit pressure and density

Unstructured Grids

```
Q [LIMIT] [PRESSUREMIN pmin] [PRESSUREMAX pmax] \ [DENSITYMIN dmin] [DENSITYMAX dmax]
```

This keyword may be used, with unstructured grids only, to set limits on the pressure and density to aid convergence. The limits pmin, pmax, dmin, and dmax are non-dimensional, normalized by the freestream static values. The default values are pmin = dmin = 0.01 and pmax = dmax = 250. In practice, setting the maximums to about two or three times the stagnation values is reasonable.

Note while the limits can be set to very small (or large) values, the limiting cannot be turned off. Specifying a limit as zero or a negative value results in the default being used.

See Also: DQ LIMITER, FIXER

REINITIALIZE — Reinitialize selected flowfield zones on restart

```
REINITIALIZE [MODE SOLUTION] [ZONE] range1[, range2[, ... rangen]] [IJK_RANGE]
```

This keyword may be used to reinitialize the flow conditions after a restart. The reinitialization will be done as if Wind-US were being run from scratch, i.e., to freestream values, or (for structured zones) to values specified using the ARBITRARY INFLOW and/or BL_INIT keywords, as described in Section 3.6 starting on p. 39.

In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

For structured zones, the IJK_RANGE option may be used in conjunction with the ARBITRARY INFLOW keyword to reinitialize only selected portions of the flow. Conditions will be reinitialized only at those points specified via the IJK_RANGE parameter on the ARBITRARY INFLOW keyword. If the IJK_RANGE parameter is not specified with the REINITIALIZE keyword, the conditions at all the grid points in the specified zones will be reinitialized.

Multiple REINITIALIZE keywords are permitted, each on a separate line in the input data file.

When the REINITIALIZE keyword is not used, a restart run will abort if the grid size for a zone in the grid (.cgd) file is different from its size in the flow (.cfl) file (unless ITERATIONS PER CYCLE is set to -2 for the zone in question, indicating that it plays no part in the calculation). However, when the REINITIALIZE keyword is used,

- The solution will be reinitialized in the specified zones, and if the grid sizes in the .cgd and .cfl files don't match, the size in the .cfl file will be reset to match the size in the .cgd file.
- In addition, even for zones not explicitly specified, if the grid sizes in the .cgd and .cfl files don't match, the solution will automatically be reinitialized, and the size in the .cfl file will be reset to match the size in the .cgd file.

Example

The following example will reinitialize flow conditions in zones 1 and 3. If the grid sizes for those zones aren't the same in the existing .cgd and .cfl files, the grid size in the .cfl file will be reset to match the size in the .cgd file. In addition, if the grid size for any other zone is different in the .cgd and .cfl files, that zone will also be reinitialized, and the grid size in the .cfl file will be reset to match the size in the .cgd file.

```
REINITIALIZE ZONE 1
REINITIALIZE ZONE 3
```

See Also: ARBITRARY INFLOW, BL_INIT, FREESTREAM

RELAX COUPLING — Set zone coupling relaxation factor

Structured Grids

```
RELAX {COUPLE | COUPLING} factor \
[ZONE range1[,range2[, ... rangen]] \
[[BOUNDARY] {ALL | I1 | IMAX | J1 | JMAX | K1 | KMAX}]]
```

This keyword allows the user to specify the relaxation factor when using characteristic zone coupling and structured grids. *factor* may range from 0.0 for no coupling, to 1.0 for full zone coupling. In the zone specification, the *range* parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

begzone: Selects all zones from begzone to MAXZONE, the total number of zones in the

grid file

: endzone Selects all zones from 1 to endzone

ALL Selects all zones

If the zone specification is omitted, the specified *factor* will be applied at all boundaries in all zones. Note that a boundary cannot be specified without the zone specification.

If a zone (or zones) is specified, but the boundary is not (or the boundary is specified as BOUNDARY ALL), the specified factor will be applied at all boundaries in the specified zone(s).

The default relaxation factor is 0.7 for steady-state calculations, and 1.0 for for space-marching and time-accurate calculations. If the RELAX COUPLING keyword is used, and all zones are selected (either by omitting the zone specification, or by specifying ZONE ALL), the relaxation factor is automatically set to 1.0 for space-marching and time-accurate calculations. However, if specific zones were selected, the specified *factor* will be used at the specified boundary in those zones, even for space-marching and time-accurate calculations.

REL-ROT-ZONE — Relative rotating zones (block)

Structured Grids

```
REL-ROT-ZONE

ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX}

ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX}

ROTATING-ZONE AVERAGE [ABOUT] {X|Y|Z}

ENDRRZ
```

For structured grids, the REL-ROT-ZONE keyword block, along with the ROTATE keyword, may be used to specify that one zone is rotating relative to another zone. This "relative rotating zone" capability is intended to simulate rotating devices such as compressor fans. The ROTATE keyword is used to specify which zone(s) are rotating, plus the center of rotation and the rotation rate. The REL-ROT-ZONE keyword block specifies the location of the interface between the two zones, and how flow conditions are to be transferred between zones.

The axis of rotation must align with one of the Cartesian coordinate axes. The interface between the zones must correspond to a surface of revolution and grid lines in the circumferential direction must be at a constant radius relative to the rotation axis.

The elements of the REL-ROT-ZONE keyword block are defined as follows:

```
REL-ROT-ZONE
...
ENDRRZ
```

The REL-ROT-ZONE and ENDRRZ keywords define the beginning and end of the REL-ROT-ZONE block.

```
ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX}
```

These two lines define the interface between the two zones. The relevant zones are given by the values of iz1 and iz2, and the relevant boundaries within zones iz1 and iz2 are specified via the BOUNDARY keyword parameter.

- iz1 Zone to which increments will be added when passing information to iz2
- iz2 Zone receiving positive increments, increments will be subtracted when passing information back to zone iz1

```
ROTATING-ZONE AVERAGE [ABOUT] {X|Y|Z}
```

When the relative-rotating-zone capability is used, flow conditions at each radial grid point are circumferentially averaged before sending them to the adjacent coupled zone. This permits the communication of the bulk fluid properties between zones, while maintaining radial distributions and the efficiency of local time stepping. The ROTATING-ZONE AVERAGE keyword defines the circumferential direction used for the averaging.

Note that since it's currently assumed that the axis of rotation aligns with one of the Cartesian coordinate axes, the circumferential direction specified with ROTATING-ZONE AVERAGE must be consistent with the rotation rate specified with with the ROTATE keyword.

 $See \ Also: \ {\tt ROTATE}$

RESIDUAL_OUTPUT — Residual output controls

```
RESIDUAL_OUTPUT {MAX [NUM num] [FREQUENCY iterfreq] | \ STATISTICS [{LINEAR|LOGARITHMIC} nbins] [FREQUENCY iterfreq]}
```

This keyword may be used to write residual data for individual equations to the list output (.lis) file. There are two basic options, MAX and STATISTICS.

If RESIDUAL_OUTPUT MAX is specified, residuals for each equation will be written every *iterfreq* iterations at *num* individual grid points, in descending order of the absolute value of the residual. The defaults for *iterfreq* and *num* are 50 and 100, respectively. For structured grids the form of the output for each grid point is

```
*MAXNS*
                iter zone eq i j k res
and for unstructured grids the form is
     *MAXNS*
                iter zone eq cell res
where
   iter
           Iteration number
           Zone number
   zone
           Equation number (i.e., 1–5 corresponding to the continuity, x-, y-, and z-momentum,
   eq
           and energy)
           Grid point indices
   i, j, k
   cell
           Cell number
```

If RESIDUAL_OUTPUT STATISTICS is specified, residual statistics for each equation will be written every *iterfreq* iterations. The range of residuals from the minimum to the maximum value will be divided into *nbins* "bins," distributed either linearly or logarithmically between the two values. The default for *iterfreq* is 50, and the default number of bins is 10, distributed linearly. The form of the output for each bin is

STATNS iter zone eq res_ctr res_n res_%

```
where

iter Iteration number

zone Zone number

eq Equation number (i.e., 1–5 corresponding to the continuity, x-, y-, and z-momentum, and energy)

res_ctr Residual value at center of bin

res_n Number of grid points with residuals in the bin
```

The RESIDUAL_OUTPUT keyword may be specified twice to write both residuals and statistical data to the .lis file.

Percentage of the total number of grid points with residuals in the bin

Residual

res

 $res_{-}\%$

RESTART | START — Begin run in specified zone

{START | RESTART} [zone [plane]]

This keyword allows the user to specify the zone (and plane, for a marching solution) in which the solution will start (or restart) when running in single-processor mode. In parallel mode, the starting zone number is automatically set to 1.

In both single-processor and parallel modes, if this keyword is not used, Wind-US checks for the existence of a flow (.cfl) file. If one exists, the solution will be restarted using the values in the .cfl file as initial conditions. If a .cfl file does not exist, Wind-US will create and initialize one.

zone First zone entered. If not specified, the solution will begin in the zone after the last successfully completed zone. However, if Global Newton iteration is being used, the solution must start in zone 1, and specification of a zone value other than 1 is not allowed.

plane i-plane to start/restart solution when running in PNS marching mode. If omitted, Wind-US will automatically start at the next i-plane after the last one completed.

RHS — Explicit operator control

Structured Grids

```
RHS scheme [order [modifier]] [ZONE range1[,range2[, ... rangen]]]
```

Unstructured Grids

```
RHS scheme [order] [ZONE range1[,range2[, ... rangen]]]
```

This keyword allows control of the explicit operator used within each zone. The parameter *scheme* specifies the general type of differencing scheme, *order* specifies the differencing order, and (for structured grids) *modifier* specifies the type of spatial differencing.

The zone specification format is essentially the same as for a *zone_selector* (see Section 10.2), except that the word ZONE is required if a *range* is being input.

Structured Grids

For structured grids, the following choices are available for *scheme*:

```
CENTRAL, COAKLEY, HLLC, HLLE, ROE, ROE_OVER, RUSANOV, VANLEER
```

The parameter *order* specifies the differencing order, and must be one of the following:

```
FIRST, SECOND, THIRD, FOURTH, FIFTH
```

The *modifier* must be one of:

```
CENTRAL, UPWIND, PHYSICAL, UPWINDBIASED
```

Not all combinations of options are valid.

If *scheme* is CENTRAL, second-order central differencing is used, and any values specified for *order* and *modifier* are ignored.

If *scheme* is COAKLEY, Coakley differencing is used and the following options are available for *order*. Any *modifier* that is specified is ignored. Following the usual conventions for displaying keyword syntax, optional keyword parameters are inside square brackets. Thus, RHS COAKLEY and RHS COAKLEY SECOND both give second-order Coakley differencing, fully upwind.

orderExplicit OperatorFIRSTFirst-order, upwind[SECOND]Second-order, fully upwindTHIRDThird-order, upwind-biased

If *scheme* is ROE, VANLEER, HLLE, HLLC, or RUSANOV, then Roe, Van Leer, HLLE, HLLC, or Rusanov differencing is used, respectively.

If *scheme* is ROE_OVER, an alternative implementation of the Roe scheme from the OVERFLOW code is used. This implementation seems to be faster, and includes an entropy fix that prevents expansion shocks.

The HLLE scheme also includes a built-in entropy fix to prevent expansion shocks. Otherwise, the HLLE and Roe schemes give very similar results.

The following options are available for *order* and *modifier* with all of the Roe, Van Leer, HLLE, HLLC, and Rusanov schemes. Again, optional parameters are inside square brackets. Thus, RHS ROE, RHS ROE SECOND, and RHS ROE SECOND PHYSICAL all give second-order Roe upwind-biased differencing, modified for stretched grids.

order and modifier Explicit Operator FIRST [UPWIND] First-order, upwind FIRST PHYSICAL First-order, upwind, modified for stretched grids [SECOND [PHYSICAL]] Second-order, upwind-biased, modified for stretched grids SECOND CENTRAL Second-order, central SECOND UPWINDBIASED Second-order, upwind-biased SECOND UPWIND Second-order, fully upwind THIRD [UPWINDBIASED] Third-order, upwind-biased THIRD UPWIND Third-order, fully upwind FOURTH [UPWINDBIASED] Fourth-order, upwind-biased FOURTH CENTRAL Fourth-order, central FIFTH [UPWINDBIASED] Fifth-order, upwind-biased

If the RHS keyword is not used, the second-order upwind-biased Roe scheme with modifications for stretched grids (i.e., RHS ROE SECOND PHYSICAL) is used.

Notes

- Roe zonal coupling requires that Roe, Van Leer, HLLE, HLLC, or Rusanov differencing be used.
- TVD flux limiting, and the default zonal coupling procedure (high-order Roe), cannot be used with some of the higher-order explicit operators. See the TVD and COUPLING keywords for details.
- Some of the explicit operators cannot be used in a spatial marching calculation. See the MARCHING keyword for details.
- If a first-order upwind explicit operator modified for stretched grids is used, test option 189
 must be set.
- A Roe, Van Leer, HLLE, HLLC, or Rusanov explicit operator must be used in zones containing holes.
- The ROE_OVER scheme cannot currently be used for multi-species flows.
- The Van Leer scheme cannot be used with ROTATE or GRAVITY.

See Also: COUPLING, MARCHING, TVD, HLLE, ENTROPY FIX, TEST 189

Unstructured Grids

For unstructured grids, the following choices are available for *scheme*:

```
HLLC, HLLE, ROE, RUSANOV
```

The parameter order specifies the differencing order, and must be one of the following:

```
FIRST, SECOND
```

The default for order is SECOND.

If the RHS keyword is not used, the second-order HLLE scheme (i.e., HLLE SECOND) is used.

```
See Also: ENTROPY FIX
```

ROLL — Specify roll about one of the coordinate axes

ROLL {X | Y | Z} rat x1 x2

The ROLL keyword allows the user to specify roll about one of the coordinate axes.

rat Rotation rate in degrees per second. (Note: prior to Wind-US 2.184 this was in radians per second. Input data files using this keyword with earlier versions of Wind-US will need to be changed if used with Wind-US 2.184 or later.)

x1, x2 Coordinates of the center of rotation. The coordinates specified by x1 and x2 depend on the rotation axis, as shown in the following table:

Rotation axis	x1	x2
X	\overline{y}	z
Y	z	x
Z	x	y

Note that in this mode, the calculation is done in inertial space, and the grid is assumed to roll through inertial space at the specified rate. The grid velocity imparts the rotating component. Thus, in order to show the velocity relative to the rotating axes, the roll component of velocity must be subtracted (via the CFPOST post-processing package or another mechanism).

Note also that, unfortunately, unless ROLL is used in a true time-accurate mode which moves the grid each time step (not currently available), as well as giving it velocity, the ROLL capability neglects the rotational potential energy, and so is not correct for high rotation rates.

With this keyword, the velocity vector must be aligned with the coordinate axis. (To get an angle of attack, rotate the grid in GMAN.)

 $See \ Also: \ {\tt ROTATE}$

ROTATE — Perform calculation in a rotating frame of reference

```
ROTATE x_{cntr} y_{cntr} z_{cntr} \Omega_x \Omega_y \Omega_z [zone_selector]
```

The ROTATE keyword allows the user to compute the flow in a rotating reference frame. The center of rotation and rotation rates are specified via the arguments:

 $x_{cntr}, y_{cntr}, z_{cntr}$ Center of rotation, in grid units.

 Ω_x , Ω_y , Ω_z Rotation rates about the three Cartesian coordinate axes, in degrees/sec. (Note: prior to Wind-US 2.184 these were in radians per second. Input data files using this keyword with earlier versions of Wind-US will need to be changed if used with Wind-US 2.184 or later.)

If the zone_selector is used (e.g., to specify that only selected zones are rotating), the REL-ROT-ZONE keyword block must be used to specify the location of the interface between the zones, and how flow conditions are to be transferred between zones. In addition, the axis of rotation must align with one of the Cartesian coordinate axes. Thus, although all three of the rotation rates Ω_x , Ω_y , and Ω_z must be specified, two of them must be zero. The use of zone_selector with the ROTATE keyword is only available for structured grids.

Note that a similar capability is given by the ROLL keyword, in which the grid is rotated about an axis. However, ROLL does the calculation in inertial space, and uses the grid velocity to impart a rotating component. Unfortunately, unless ROLL is run in a true time-accurate mode which moves the grid each time step (not currently available), as well as giving it velocity, the ROLL capability neglects the rotational potential energy, and so is not correct for high rotation rates.

This keyword cannot be used with the Van Leer differencing scheme (RHS VANLEER).

See Also: REL-ROT-ZONE, ROLL, ARBITRARY INFLOW

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SCHMIDT — Set the laminar and turbulent Schmidt numbers

SCHMIDT Schl Scht

This keyword may be used to set the laminar and turbulent Schmidt numbers. The default values are 0.72 and 0.9, respectively.

${\bf SEQUENCE-Grid\ sequencing\ control}$

Structured Grids

SEQUENCE iseq jseq kseq [zone_selector]

For structured grids, this option allows the user to, in effect, thin the computational grid, thus reducing CPU requirements. It can be used to obtain quick starting solutions for the fine grid. Separate keywords may be used for each zone.

iseq Number of sequencing levels in the i direction

jseq Number of sequencing levels in the j direction

kseq Number of sequencing levels in the k direction

For one sequencing level, every other grid point is used in the calculation. For two sequencing levels, the sequencing process is repeated, resulting in only every fourth grid point being used. And so on. At the end of the run, the solution is interpolated back onto the entire grid to aid in restarting the solution and to provide a continuous field for post-processing.

There are no restrictions on the number of points in the sequenced direction, except that there must be at least five points in the sequenced (i.e., coarse) grid.

SMOOTHING — Add dissipation to explicit operator

Structured Grids

```
SMOOTHING [SECOND val1] [FOURTH val2] [SMLIMT val3] [zone\_selector]
```

This keyword is used to specify artificial dissipation to be added to the explicit operator. The values *val1*, *val2*, and *val3* are the second-order smoothing coefficient, the fourth-order smoothing coefficient, and the smoothing limiter, respectively.

If the SMOOTHING keyword isn't used, no smoothing is applied. If the SMOOTHING keyword is used, but without specifying values, the default values listed below are used in the selected zones. No smoothing is done in non-selected zones. (The table also includes a "TVD-like" value for each smoothing parameter.)

Parameter	Default	"TVD-Like" Value
val1	1/4	< 1
val2	1/256	< 1/32
val 3	0	1

The default values shown in the above table represent a Jameson-type smoothing, and are good for an Euler analysis. For viscous problems, it is recommended that val1, val2, and val3 be set equal to 1, 1/64, and 2, respectively.

See Also: BOUNDARY-DAMP, ACCELERATE, TEST 49

SOLVER-STAGES — Staged solution input (block)

```
SOLVER-STAGES
   [RESTART_STAGES]
   [STEP stage_name
      SOLVER [solver_name]
          [SET SOLVER_STEPS [TO] nSteps]
          [SET TIME_LEVELS [TO] ntlvls]
          [SET CYCLES [TO] numcyc]
          [SET ITERATIONS [PER] [CYCLE] | ITER_CYCLE [TO] niters [zone_selector]]
          [SET CFL [T0] cfl1 [zone_selector]]
          [SET TVD [T0] factor [zone_selector]]
          [SET ALPHA [TO] alpha]
          [SET BETA [TO] beta]
          [SET SEQUENCE [TO] iseq jseq kseq [zone_selector]]
          [SET TURBULENCE [TO] EULER | LAMINAR [zone_selector]]
          [SET IMPLICIT [TO] NONE [zone_selector]]
          [SET RHS [TO] scheme [order [modifier]] \
             [ZONE range1[, range2[, ... rangen]]]
      ENDSOLVER
   ENDSTEP]
ENDSOLVER-STAGES
```

This keyword block is used to define input for a staged solution. In a staged solution, changes can be made to various input options that would otherwise require a stopping and restarting the solution.

The various elements of the SOLVER-STAGES keyword block are defined as follows:

```
SOLVER-STAGES
...
ENDSOLVER-STAGES
```

The SOLVER-STAGES and ENDSOLVER-STAGES keywords define the beginning and end of the SOLVER-STAGES block.

```
RESTART_STAGES
```

This keyword should be used when restarting a staged solution. The solution will continue with the stage in effect at the end of the previous run.

```
STEP stage_name
...
ENDSTEP
```

The STEP/ENDSTEP keywords bracket input for a single stage, where $stage_name$ is the user-specified name for the stage. Multiple stages are defined by using multiple STEP/ENDSTEP sections.

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```
SOLVER [solver_name] ... ENDSOLVER
```

The SOLVER/ENDSOLVER keywords bracket input for a particular solver in the current stage, where *solver_name* is the name of the solver to be used. The default, and only valid solver name in Wind-US, is NAVIER-STOKES. Note that the SOLVER/ENDSOLVER keywords must be specified.

```
SET SOLVER_STEPS [TO] nSteps
```

Specifies the number of times the current solver should be run in the current stage. The default is 1.

```
SET TIME_LEVELS [TO] ntlvls
```

Specifies the number of global Newton time levels to advance. The default is the value specified with the NEWTON keyword, or in the TEMPORAL keyword block, whichever is used to specify Newton iteration. If Newton iteration is not being used, this keyword should not be specified, and *ntlvls* will be automatically set to 1.

```
SET CYCLES [TO] numcyc
```

Specifies the number of cycles to run. The default is the value specified with the CYCLES keyword.

```
SET ITERATIONS [PER] [CYCLE] | ITER_CYCLE [TO] niters [zone_selector]
```

Specifies the number of iterations per cycle. The default is the value specified with the ITERATIONS keyword.

```
SET CFL [T0] cfl1 [zone_selector]
```

Specifies the CFL number, time step, or time step parameter to be used. The default is the value specified with the CFL keyword. Note that the options and values specified with the CFL keyword, in particular the MODE option and the choice between CFL and SECONDS, are still relevant for the current stage, even if a different value is specified here for cfl1. Note also that if MODE 2 is used with the CFL keyword, it should only be used at the start of the run, and SET CFL shouldn't be specified in the SOLVER-STAGES block for the first stage.

```
SET TVD [T0] factor [zone_selector]
```

Specifies the TVD factor to be used. The default is the value specified with the TVD keyword. Note that the type of TVD limiter (for structured grids) or slope limiter (for unstructured grids) is controlled by the options used with the TVD keyword, even if a different value is specified here for factor.

```
SET ALPHA [TO] alpha SET BETA [TO] beta
```

Specifies the freestream angles of attack and yaw, in degrees. The defaults are the values specified with the FREESTREAM keyword.

```
SET SEQUENCE [TO] iseq jseq kseq [zone_selector]
```

Specifies the number of grid sequencing levels to be used in the i, j, and kdirections. The defaults are the values specified with the SEQUENCE keyword. This keyword is only applicable to structured grids.

```
SET TURBULENCE [TO] EULER | LAMINAR [zone_selector]
```

The SET TURBULENCE keyword may be used to specify Euler (i.e., inviscid) or laminar flow. It cannot currently be used to specify a particular turbulence model. For turbulent flow, the model to be used must be specified using the TURBULENCE keyword.

```
SET IMPLICIT [TO] NONE [zone_selector]
```

Turns off the implicit operator (i.e., switches to an explicit scheme). [Currently, if SET IMPLICIT TO NONE is specified for any zone, the implicit operator is turned off in all zones.]

```
SET RHS [TO] scheme [order [modifier]] [ZONE range1[,range2[, ... rangen]]]
```

Specifies the explicit operator to be used. The choices for *scheme*, *order*, and *modifier* are described in the RHS keyword documentation. Note that *modifier* only applies to structured grids. The default is the operator specified with the RHS keyword.

SPAWN — Run external processes from Wind-US

```
\verb|SPAWN "|process"| [FREQUENCY | freq] [CYCLES | TIME-LEVELS | STAGES] [NOCHECKPOINT]|\\
```

This keyword allows the user to start (spawn) a process without stopping Wind-US. A process can be any valid Unix script or executable. When a new process is spawned, Wind-US will wait until the spawned process completes before continuing.

Spawned processes will be run in the same directory as Wind-US. For convenience, you may thus want to use the -runinplace option with the *wind* script. See the description of the -runinplace and -runroot options to the *wind* script in Section 8.1 for details about the Wind-US run directory.

The keyword arguments and options are:

process Full path name to the process (must be in quotes, 80 characters max)

freq

Frequency for spawning the process, in terms of cycles, time levels, or stages, as determined by the option CYCLES, TIME-LEVELS, or STAGES. The default is to spawn the process every time level when Newton iteration is used, and every cycle otherwise. CYCLES may not be specified with Newton iteration unless NOCHECKPOINT is also specified.

Wind-US will tack the current cycle number (even when the frequency is in terms of time levels or stages) onto the end of the *process* call for potential use by the process as a parameter.

NOCHECKPOINT

By default, Wind-US will update the common flow (.cfl) file before spawning the process. NOCHECKPOINT will disable this update. In either case, the checkpoint interval specified by the checkpoint directive in the multi-processing control (.mpc) file will be unchanged. (See Section 9.6.)

Multiple processes may be spawned by using multiple SPAWN keywords. The FREQUENCY and NOCHECKPOINT options may be specified independently for each process.

Note that while a *process* is being executed, Wind-US's input and output files (i.e., the *.lis* file, *.cfl* file, etc.) remain connected to the Fortran units used by Wind-US. Thus, if a *process* runs a program like *resplt* or *cfpost*, these files cannot be used directly as input. Instead, the *process* must copy the needed file(s) to temporary files, and use the temporary files as input.

Example

A common use for the SPAWN keyword, especially for unsteady flows, is to save intermediate .cfl files during a Wind-US run. A shell script may be SPAWN'ed at specified intervals to save the file under a unique name.

For example, consider the following shell script, named save_cfl.

```
#! /bin/sh
#
# save_cfl - Save intermediate .cfl files. This script is intended
# to be spawned during a run using the SPAWN keyword, as
#
# SPAWN "path-to-script path-to-cfl" FREQUENCY num
#
# where 'path-to-script' is the full path name for this
```

```
#
             script, 'path-to-cfl' is the full path name for the saved
             .cfl files (the current cycle number will automatically
             be appended to this name), and 'num' is the frequency for
             the SPAWN command. Note that the quote marks around
             "path-to-script path-to-cfl" must be included.
# Check for enough arguments
if [ $# -1t 2 ]
then
   echo 'Error: save_cfl: Path name for saved .cfl files not specified' >& 2
   echo '
               Use SPAWN keyword, as: ' >& 2
   echo '
               SPAWN "path-to-script path-to-cfl" FREQUENCY num' >& 2
   exit 1
fi
# $1 is path name for the saved .cfl files, $2 is current cycle number
cp FORO20 $1.$2
```

If the above script is located in the directory /home/user/bin, the SPAWN keyword may be used as follows to save the .cfl file every 1000 cycles (assuming a non-Newton run) in the directory /home/user/wind.

```
SPAWN "/home/user/bin/save_cfl /home/user/wind/runa.cfl" FREQUENCY 1000
```

Since the cycle number is automatically added to the end of the *process* when it's spawned, the above use of the SPAWN keyword causes the *save_cfl* script to be run as

```
/home/user/bin/save_cfl /home/user/wind/runa.cfl ncycle
```

where ncycle is the current cycle number. The result is that the intermediate .cfl files will be stored in the directory /home/user/wind, with names runa.cfl.1000, runa.cfl.2000, etc.

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STAGES — Multi-stage time stepping

Structured Grids

STAGES
$$ityp$$
 [COEFFICIENTS $x1$ $x2$... xn] [$zone_selector$]

For structured grids, this keyword allows the user to use a Runge-Kutta time stepping option. The parameter ityp specifies the number of stages, and $x1 \ x2 \dots xn$ specify the coefficients. The following ityp values are available:

ityp Stages

- 0 One stage, Euler implicit or explicit
- 1 Four stage, Jameson type
- Three stage, minimal storage, third order

The default coefficients are:

\underline{ityp}	# Req'd	<u>Defaults</u>
0	_	_
1	4	1/4, 1/3, 1/2, 1
5	5	1/4, 8/15, 0, 5/12, 3/4

$ext{TDA_INVALID}$ — Flag invalid boundary data in .tda file

TDA_INVALID [ZONE] range1[,range2[, ... rangen]]

This keyword tells Wind-US that the boundary data from the .tda file for the specified zone(s) should not be used, until new boundary data is written to the file during the first solution cycle. In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

See Also: TDA_VALID

TDA_VALID — Flag valid boundary data in .tda file

```
TDA_VALID [ZONE] range1[,range2[, ... rangen]]
```

This keyword tells Wind-US that the boundary data in the .tda file for the specified zone(s) may be used, and doesn't need to be checked for changes in run conditions, sizes, etc. Zones already marked as invalid with TDA_INVALID are left as is, not changed to valid. In the zone specification, the range parameter(s) must be one of the following forms:

zonenum Selects zone zonenum

begzone: endzone Selects all zones from begzone to endzone

ALL Selects all zones

See Also: TDA_INVALID

TEMPORAL — Time-marching input (block)

```
TEMPORAL

[TIME LEVELS ntlvls

NEWTON [[PHYSICAL] TIMESTEP NewtonDt]

[CONVERGENCE {LEVEL|ORDER} newcvg]]

[IMPLICIT TIME {FIRST|SECOND} [ORDER] [ZONE range1[,range2[, ... rangen]]]]

[CFTVARS {LOCAL|LINK}]

[NEWTVARS {LOCAL|LINK}]

[SORDTVARS {LOCAL|LINK}]

[SAVE {NEWTON | NO_NEWTON}]

[SAVE {TIME | NO_TIME}]

ENDTEMPORAL
```

The TEMPORAL keyword block is used for various input options related to the time-marching procedure to be used. The various elements of the TEMPORAL keyword block are defined as follows:

TEMPORAL

Defines the beginning of the TEMPORAL block

```
TIME LEVELS ntlvls

NEWTON [[PHYSICAL] TIMESTEP NewtonDt]

[CONVERGENCE {LEVEL|ORDER} newcvg]]
```

The TIME LEVELS keyword may be used to specify that the Global Newton time stepping algorithm is to be used. The parameter ntlvls specifies the number of Global Newton time levels to advance, and must be specified. The value NewtonDt is the size of the physical Newton time step, in seconds. If not specified, the minimum zonal value from the CFL keyword is used.

There are two possible procedures for determining the overall global convergence with Global Newton iteration.

CONVERGENCE LEVEL Convergence is assumed when

$$|\mathbf{Q}^n - \mathbf{Q}^{n-1}| < newcvg$$

where \mathbf{Q} represents the vector of dependent variables, and n is the Newton time level.

CONVERGENCE ORDER Convergence is assumed when

$$\frac{|\mathbf{Q}^n - \mathbf{Q}^{n-1}|}{\max(|\mathbf{Q}^n - \mathbf{Q}^{n-1}|)} < 10^{-newcvg}$$

The default, if the CONVERGENCE keyword isn't used, is CONVERGENCE ORDER 3.

Within a Newton time level, the number of cycles and iterations to be run is specified using the CYCLES and ITERATIONS keywords.

The convergence criteria within a Newton time level may be specified using the CONVERGE keyword. Note that for structured grids the default for the CONVERGE keyword is a four-order of magnitude decrease in the maximum residual. If Newton iteration is being used for a steady flow problem

with structured grids, with the default of three orders of magnitude for the global convergence criteria, it would make sense to also use three orders of magnitude for the convergence criteria within a Newton time step.

See Also: CFL, CYCLES, ITERATIONS, CONVERGE, NEWTON

```
IMPLICIT TIME {FIRST|SECOND} [ORDER] [ZONE range1[, range2[, ... rangen]]]
```

This keyword sets the order of the implicit time marching scheme. The default, if the IMPLICIT TIME keyword isn't used, is first order.

Second-order time marching may be used with both structured and unstructured grids, and with or without Global Newton time marching. With structured grids, it may be used with any implicit operator except MacCormack's modified approximate factorization procedure (IMPLICIT MACCORMACK).

The zone specification format is essentially the same as for a *zone_selector* (see Section 10.2), except that the word ZONE is required if a *range* is being input.

See Also: IMPLICIT, IMPLICIT ORDER

CFTVARS {LOCAL|LINK}
NEWTVARS {LOCAL|LINK}
SORDTVARS {LOCAL|LINK}
SAVE {NEWTON | NO_NEWTON}
SAVE {TIME | NO_TIME}

Using Global Newton iteration and/or second-order time differencing requires the storage of flow field data at additional time levels. When CFTVARS LOCAL is specified, the extra time levels required for both Global Newton iteration and second-order time differencing are stored in the common flow (.cfl) file. Specifying CFTVARS LINK causes the extra time levels to be stored in a separate time data (.cft) file, and linked to the .cfl file. The NEWTVARS and SORDTVARS keywords may be used instead of CFTVARS, and apply independently to Global Newton iteration and second-order time differencing, respectively.

The default is to store the extra time levels for Newton iteration in the .cft file, and for second-order time differencing in the .cft file.

The SAVE keyword may be used to specify whether or not the data at these extra time levels is retained in the .cfl or .cft file at the end of a run. The default is to save the data for second-order time differencing, but not for Global Newton iteration.

The CFTVARS, NEWTVARS, SORDTVARS, and SAVE keywords may also be specified outside the TEMPORAL keyword block.

ENDTEMPORAL

Defines the end of the TEMPORAL block.

TEST — Non-production test options

TEST number [mode]

For purposes of program testing, the user may select one or more test options for any job. These options are selected by the keyword TEST (beginning in column 1) followed by one or two integer variables. The keyword and the integer variables must be separated by one or more blanks. The first integer selects the test option. The second integer, which is not mandatory, selects a mode for that option. If the second integer is omitted, a default mode is activated. Any number of valid test options may be selected. However, only one mode may be activated for each option. If an option is selected more than once, the mode identified in the last selection will be used.

The list of valid options and modes is provided in Section 11.

TSL | THIN SHEAR LAYER — Thin shear layer option

Structured Grids

{TSL | THIN} [SHEAR] [LAYER] word1 word2 word3 [zone_selector]

For structured grids, this keyword turns off the viscous terms in specified coordinate directions. The parameters word1, word2, and word3 are keywords controlling the viscous terms in the ξ -, η -, and ζ -directions, respectively. They may have the following values:

 Keyword
 Meaning

 VISCOUS
 Retain all viscous terms in this direction

 INVISCID
 Neglect all viscous terms in this direction

By default, when INVISCID is set for a direction, the scalar implicit operator will be used in that direction (the full block implicit operator is used in viscous directions). This may be overridden using the IMPLICIT keyword.

TTSPEC — Wall temperature and transition (block)

Structured Grids

```
TTSPEC

TYPE {TEMPERATURE | TRANSITION}

ZONE zone_number

MODE {CURRENT | FILE filename | CHT}

UPDATE [INTERVAL] frequency

SURFACE surfname

ENDTTSPEC
```

For structured grids, this keyword block allows a point-by-point surface temperature boundary condition to be used, and/or point-by-point boundary layer transition information to be specified. Surface temperature and transition data may be specified for any boundary surface in any zone.³⁶ The maximum total number of temperature/transition surfaces, combined, is determined by the Fortran parameter MAXTTS, currently equal to 20.

The normal procedure is to do an initial run, perhaps with just a few cycles, without specifying temperature/transition information, to create the initial common flow (.cfl) file. The temperature/transition data is then created and added to the .cfl file using the tmptrn utility.

For backward compatibility with early versions of WIND, however, the temperature/transition data may be read from separate ASCII files, one for each applicable surface, created with the older *tmptrn* utility. This would only be necessary during the initial (i.e., non-restart) run, since Wind-US will automatically store the data in the .cfl file, where it can be read during subsequent runs.

In addition, in earlier versions of *tmptrn*, the temperature written into the *.cfl* file was in K, instead of being non-dimensionalized. This error was fixed in version 1.8 of *tmptrn*. However, some versions of Wind-US had coding to accommodate the dimensional temperature value, and these versions will not work with temperature distributions written using *tmptrn* 1.8 and above. See the *tmptrn* documentation for details.

The various elements of the TTSPEC keyword block are defined as follows:

```
TYPE {TEMPERATURE | TRANSITION}
```

This keyword indicates whether a surface temperature distribution or boundary layer transition information is being specified.

```
{\tt ZONE} \ \ zone\_number
```

This keyword specifies the number of the zone containing the boundary surface along which temperature or transition data is being specified.

```
MODE {CURRENT | FILE filename | CHT}
```

This keyword specifies where the surface temperature or transition data is obtained.

 $^{^{36}}$ Note, though, that although Wind-US allows temperature/transition data to be specified on any boundary surface, the *tmptrn* utility used to add the data to the *.cft* file currently does not allow specification on the i_1 or i_{max} surface.

CURRENT The data is stored in the flow (.cfl) file.

FILE filename The data is stored in the ASCII file filename, created using the older tmptrn

utility distributed with early versions of WIND. If filename is specified without

an extension, an extension of .dat is assumed.

This option is included for backward compatibility only, using existing files created with the older *tmptrn* utility, and is only needed for an initial (i.e., non-restart) run. Subsequent (i.e., restart) runs should use MODE CURRENT. New applications should use the latest *tmptrn* utility to write the tempera-

ture/transition data into the .cfl file directly.

CHT Wall temperatures are computed using an external loosely-coupled conjugate

heat transfer code. The Wind-US executable being used must contain the appropriate Surface Data Interface library routines, not currently distributed with Wind-US. This mode only applies when temperature is being specified,

not boundary layer transition.

This mode is still under development, and thus not yet fully operational.

```
{\tt UPDATE \ [INTERVAL]} \ \mathit{frequency}
```

When MODE CHT is used, wall temperatures will be updated every *frequency* cycles. This keyword does not apply to other MODE options.

```
{\tt SURFACE}\ surfname
```

This keyword specifies the boundary surface in zone zone_number along which temperature or transition data is being specified. Currently, the applicable surface names are I1, IMAX, J1, JMAX, K1 and KMAX, corresponding to the i_1 , i_{max} , j_1 , j_{max} , k_1 and k_{max} surfaces, respectively.

Note: In versions of Wind-US prior to 3.39, the keyword SUBSET ALL had to be specified, once for each surface, to properly update the total number of temperature/transition surfaces. That's not only no longer necessary, but *should not* be done. Existing input data (.dat) files should be modified to remove any SUBSET ALL keywords in TTSPEC keyword blocks.

Example

As an example, suppose we are specifying the wall temperature distribution along the j_1 surface in zone 1, and the j_1 and j_{max} surfaces in zone 2. In addition, suppose we're specifying boundary layer transition along the j_1 surface in zone 1. If existing temperature/transition files are being used, created with the older tmptrn utility, the TTSPEC keyword block for the initial run might be

TTSPEC

```
TYPE TEMPERATURE

ZONE 1

MODE FILE temp.z1j1.asc
SURFACE J1

ZONE 2

MODE FILE temp.z2j1.asc
SURFACE J1

MODE FILE temp.z2jm.asc
SURFACE JMAX

TYPE TRANSITION
```

```
ZONE 1

MODE FILE tran.z1j1.asc
SURFACE J1
ENDTTSPEC
```

Since the temperature/transition data will be written into the .cfl file during the initial run, later (i.e., restart) runs could use

```
TTSPEC

TYPE TEMPERATURE

ZONE 1

MODE CURRENT
SURFACE J1

ZONE 2

MODE CURRENT
SURFACE J1
SURFACE JMAX
TYPE TRANSITION
ZONE 1

MODE CURRENT
SURFACE J1
ENDTTSPEC
```

See Also: WALL TEMPERATURE

TURBULENCE — Turbulence model selection

Structured Grids

```
TURBULENCE [MODEL] model [ITERATIONS iter] \ [RC | ROTATION] [F2FIX dmax] [zone\_selector]
```

Unstructured Grids

```
TURBULENCE [MODEL] model [ITERATIONS iter] [zone\_selector]
```

The TURBULENCE keyword is used to request an inviscid or viscous solution, and to select a turbulence model for one or more zones. In addition to the TURBULENCE keyword itself, several additional keywords affect various aspects of the turbulence modeling procedure. Of these, see especially the MAX_WALL_DISTANCE keyword, used to specify the maximum wall distance to be used. For problems with well-behaved boundary layers, appropriate use of this keyword can significantly speed up the wall distance calculation.

- Model Selection
 - TURBULENCE Turbulence model selection
- Wall Distance
 - MAX_WALL_DISTANCE Maximum wall distance
- Spalart-Allmaras and Goldberg Keywords
 - FREE_ANUT Freestream eddy viscosity
- SST Keywords
 - COMPRESSIBLE, PRESSURE Compressibility correction options
 - FREE_K, FREE_OMEGA Freestream k and ω
 - SST COEF_PTM1 Production term modification
- k- ϵ Keywords (structured grids only)
 - K-E INITIALIZE Initialization procedure
 - K-E FREE_K, K-E FREE_MUT Freestream k and μ_t
 - K-E REINITIALIZE Reinitialize k, ϵ , and μ_t
 - K-E LINEARIZATION Linearization procedure
 - K-E TVD ORDER TVD accuracy order
 - K-E RELAX Relaxation iterations
 - K-E MAXIMUM VISCOSITY, K-E REFERENCE VELOCITY Limiting procedure
 - K-E COMPRESSIBILITY Compressibility correction options
 - K-E CMU Variable C_{μ} option
 - K-E CMUMIN Minimum C_{μ}
- Combined RANS/LES Models
 - DES Spalart DES Model
 - MDES Shih's Modified DES Model
 - LESB SST with ϵ Limiter
 - HYBRID Nichols-Nelson Hybrid Model
 - PRNS Partially Resolved Numerical Simulation (unstructured grids only)

- DETACHED-PRNS - Detached Partially Resolved Numerical Simulation (unstructured grids only)

Model Selection

The TURBULENCE keyword (see p. 260 for the syntax) is used to select inviscid, laminar, or turbulent flow, and for turbulent flow, the turbulence model to be used. The choice is determined by the input value *model*, which must be one of the following. Unless noted otherwise, these apply to both structured and unstructured grids.

{INVISCID | EULER} [NOSLIP]

Inviscid flow. If NOSLIP is specified, no-slip boundary conditions will be applied at adiabatic and viscous walls; otherwise, slip boundary conditions will be applied.

LAMINAR. Laminar flow.

CEBECI [SMITH] Cebeci-Smith algebraic turbulence model. This model is only available

for structured grids.

[BALDWIN] LOMAX Baldwin-Lomax algebraic turbulence model. This model is only avail-

able for structured grids.

PDT Combination Baldwin-Lomax and P. D. Thomas algebraic shear layer

model. This model is only available for structured grids.

[BALDWIN] BARTH Baldwin-Barth one-equation turbulence model. This model is only

available for structured grids.

SPALART [ALLMARAS] [ROUGHNESS hgt]

Spalart-Allmaras one-equation turbulence model. The ROUGHNESS option may be used to include the effects of wall roughness. The default roughness height is 0.0 (Aupoix and Spalart, 2003).

Additional keywords may be used with the Spalart-Allmaras model to specify the freestream turbulence level (p. 265), and to specify use of the Spalart or Shih's Modified Detached Eddy Simulation (DES) turbulence model (p. 271). For unstructured grids, other keywords may be used to specify use of a PRNS (Partially Resolved Numerical Simulation) or Detached PRNS model.

In multi-zone problems, the Spalart-Allmaras model may not be used with different turbulence models in other zones, other than the Spalart DES models. However, it may be used with inviscid or laminar flow in

other zones.

Goldberg pointwise one-equation turbulence model. This model is only

available for unstructured grids. An additional keyword may be used with the Goldberg model to specify the freestream turbulence level

(p. 265).

In multi-zone problems, the Goldberg pointwise model may not be used with different turbulence models in other zones. However, it may be used with inviscid or laminar flow in other zones.

POINTWISE

SST [COMPRESSIBLE | TRANSITION]

Two-equation Shear Stress Transport turbulence model. If COMPRESS-IBLE is specified, the compressibility corrections of Suzen and Hoffmann are applied. If TRANSITION is specified, a compressible version of the SST model with transition modeling is used.

Additional keywords specific to the SST model are available for specifying compressibility correction options (p. 265), freestream values of k and ω (p. 265), and the production term modification coefficient for transition modeling (p. 266). Other keywords may be used to specify use of a combined SST/LES model using an epsilon limiter (p. 271), or the Nichols-Nelson hybrid SST/LES model (p. 271). And for unstructured grids, the PRNS (Partially Resolved Numerical Simulation) model may be used.

Specifying COMPRESSIBLE has been found to be beneficial for free shear layers in flows where the convective Mach number is above about 0.5). The overall effect is to increase the spreading in mixing layers, jets, etc.

The TRANSITION option uses the compressible version of the SST model with transition modeling developed by Menter, et. al. (2003), with modifications by Denissen, Yoder, and Georgiadis (2008). This option is only available for structured grids, and is fairly sensitive to grid resolution, particularly in the streamwise direction.

Note that the COMPRESSIBLE and TRANSITION options are mutually exclusive. If both are specified, only the TRANSITION option will be used.

In multi-zone problems, the SST model may not be used with different turbulence models in other zones, other than the hybrid SST/LES model. However, it may be used with inviscid or laminar flow in other zones

CHIEN [K-E]

Low Reynolds number two-equation k- ϵ turbulence model of Chien. This model is only available for structured grids.

The user should be aware of additional keywords specific to the k- ϵ models, particularly those used to control initialization, starting on p. 266. Other keywords may be used to specify use of a Nichols-Nelson hybrid Chien/LES model, as described starting on p. 271.

In multi-zone problems, the Chien model may not be used with different turbulence models in other zones, other than the Rumsey-Gatski model and/or a hybrid Chien/LES model. It also may not be used with inviscid or laminar flow in other zones.

RUMSEY-GATSKI [ASM] [K-E]

Low Reynolds number two-equation k- ϵ algebraic Reynolds stress model of Rumsey and Gatski. This model is only available for structured grids.

The user should be aware of additional keywords specific to the k- ϵ models, particularly those used to control initialization, starting on p. 266.

In multi-zone problems, the Rumsey-Gatski model may not be used with different turbulence models in other zones, other than the Chien model. It also may not be used with inviscid or laminar flow in other zones.

Note that with the exceptions noted above for the Spalart, Goldberg, SST, Chien, and Rumsey-Gatski models, different turbulence models, or inviscid or laminar flow, may be specified in different zones. However, you must specify a "default" turbulence model (or inviscid or laminar flow) in the input data file. Wind-US will stop if you do not. By "default", we mean without specifying zones. In addition, due to a coding limitation, the zone_selector can't be used when inviscid flow is being specified.

For example, for a three-zone problem with inviscid flow in zone 1 and the Spalart-Allmaras turbulence model in zones 2 and 3, the following will *not* work because a "default" turbulence model has not been specified, and the code will stop:

```
TURBULENCE SPALART ZONE 2,3
TURBULENCE INVISCID ZONE 1
```

The following will also not work, because a zone_selector can't be used with INVISCID:

```
TURBULENCE SPALART
TURBULENCE INVISCID ZONE 1
```

Instead, one would specify the following, which will work:

```
TURBULENCE INVISCID
TURBULENCE SPALART ZONE 2,3
```

Additional options that may be specified with the TURBULENCE keyword are:

ITERATIONS iter

Wind-US organizes the equations to be solved into logical "groups" that are solved together. It also allows multiple iterations of a specific group (i.e., sub-iterations) for each "iteration per cycle". For the one- and two-equation turbulence models, the ITERATIONS option allows the user to request *iter* sub-iterations of the turbulence model equation group for each iteration per cycle. If NAVIER-STOKES ITERATIONS is defaulted, this corresponds to *iter* iterations of the turbulence model equations for each iteration of the mean flow equations.

The default for *iter* is one, indicating that each iteration per cycle corresponds to one iteration of the turbulence model equations.

RC | ROTATION

For structured grids, this option may be used with the Spalart-Allmaras one-equation model to include a correction for system rotation and streamline curvature.

F2FIX dmax

For structured grids, this option may be used with the SST two-equation model to specify a maximum distance from the wall, dmax, within which the F_2 term may be non-zero. Beyond that distance, the F_2 term is set to zero. F_2 is a blending function, designed to limit the effects of the shear stress transport term to regions near walls.

Note that for the one- and two-equation turbulence models, the turbulence model equations may be solved without simultaneously solving the Navier-Stokes equations. Of course, the turbulence variables depend on the mean flow field, so a reasonable mean flow solution must already exist.

As an example, suppose a mean flow solution has been computed using the SST turbulence model. The Chien k- ϵ variables consistent with the existing mean flow field may be computed by restarting from the SST solution, and solving just the Chien k- ϵ equations.

ITERATIONS PER CYCLE 2 ZONE ALL
NAVIER-STOKES ITERATIONS 0 ZONE ALL
TURBULENCE MODEL CHIEN ITERATIONS 5 ZONE ALL

The above keywords specify that, for all zones, there will be two iterations per cycle, with no Navier-Stokes sub-iterations and five Chien model sub-iterations for each "iteration per cycle". There will thus be a total of ten iterations of the Chien k- ϵ equations in each zone prior to completing a cycle and exchanging information between zones.

See Also: ITERATIONS, NAVIER-STOKES ITERATIONS, TEST 21, TEST 67

Maximum Wall Distance

This keyword may be used to specify the maximum distance from the wall that will be used in the algebraic models, and the Spalart-Allmaras, k- ϵ , and SST models. The value DistMax may be specified in either grid units (i.e., the same units that are used in the .cgd file) by using the GRID_UNITS modifier, or as a multiple of the "nominal" boundary layer thickness by using the BOUNDARY_LAYER_THICKNESSES (or BLT) modifier. Note that the modifier GRID_UNITS or BOUND-ARY_LAYER_THICKNESSES (or BLT) must be specified.

The "nominal" boundary layer thickness is defined as $0.37Re^{-1/5}L_c$, where L_c is a characteristic length equal to the diagonal of the bounding box containing all the viscous grid points, and the Reynolds number Re is based on L_c and the freestream flow conditions. This is empirically equal to the boundary layer thickness for incompressible turbulent flow past a flat plate at those conditions.

The default for DistMax is 10,000 grid units.

A message is periodically written to the .lis file for each zone, showing the progress of the calculation. The optional PROGRESS keyword may be used to specify how often this message is written, or to prevent it from being written at all. The value $integer_percent$ is the frequency for writing the message, expressed as a percentage of grid cells to be searched. E.g., if $integer_percent = 5$, the message will be written when the wall distance for 5% of the cells have been computed, 10% of the cells, etc. The default is to write the message, with $integer_percent$ equal to 10.

Note/Warning

The default setting for *DistMax* of 10,000 grid units is intentionally very large so that there will be essentially no chance that the cap on the distance from the wall will corrupt turbulent flow simulations. However, even though the calculation is only done once and the results are saved in the *.tda* file for later use, there may be a *very* significant penalty in computational time required at startup for flow problems with many zones and large numbers of grid points.

To significantly reduce this computational time penalty, a user should set *DistMax* to a lower value. As a general rule of thumb, if a flow problem is expected only to have attached (i.e., non-separated), relatively thin boundary layers, *DistMax* may be set to a smaller value than is needed for problems with large wakes, separated regions, or jets. In practice, it has been found that for simulations with well-behaved boundary layers, using

 ${ t MAX_WALL_DISTANCE}\ DistMax\ { t BOUNDARY_LAYER_THICKNESSES}$

where the multiplier *DistMax* is a value from 2–10, may be sufficient.

Spalart-Allmaras and Goldberg Models

 ${\tt FREE_ANUT} \ \ anut inf$

When TURBULENCE SPALART or TURBULENCE POINTWISE is specified, indicating use of the Spalart-Allmaras or Goldberg model, the separate keyword FREE_ANUT may be used to specify the freestream value of the eddy viscosity ν_t . The default value is 5.0.

SST Model

Compressibility Correction Options

If TURBULENCE SST COMPRESSIBLE is specified, indicating that the SST model is to be used with the compressibility corrections of Suzen and Hoffmann, the separate keywords COMPRESSIBLE and PRESSURE may also be specified. The COMPRESSIBLE keyword controls whether or not a compressible dissipation correction is used, and the PRESSURE keyword controls whether or not the pressure dilatation term is included. Both of these are ON by default.

Note that these keywords have no effect when TURBULENCE SST TRANSITION is specified.

Specifying Freestream k and lomega

FREE_K val_k
FREE_OM val_om

If TURBULENCE SST is specified, indicating that the SST model is to be used, the separate keywords FREE_K and FREE_OM may be used to input dimensional freestream values of k and ω . The following options are possible:

 $val_k > 0$ The turbulent kinetic energy k and the specific dissipation rate ω are specified directly, with

$$k = val k \text{ (ft}^2/\text{sec}^2)$$

 $\omega = val om \text{ (1/sec)}$

The turbulent viscosity ν_t is then equal to k/ω .

val k < 0 The turbulence intensity is set equal to |val k|, expressed as a percentage of the freestream velocity U_{∞} . Thus, the turbulent kinetic energy is computed as

$$k = 1.5(0.01 | val_k | U_{\infty})^2$$

The turbulent viscosity ν_t is automatically set equal to $0.001\nu_l$, where ν_l is the laminar viscosity, and the specific dissipation rate is computed as $\omega = k/\nu_t$.

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 $val_om < 0$ The specific dissipation rate ω is set equal to val_om percent of U_{∞}/L_{ref} , where U_{∞} is the freestream velocity, and L_{ref} is the reference length from the grid (.cgd) file. Thus

$$\omega = 0.01 \ |val_om| \ \frac{U_{\infty}}{L_{ref}}$$

The turbulent viscosity ν_t is set to the same percentage of the laminar viscosity.

$$\nu_t = 0.01 |val_om| \nu_l$$

The turbulent kinetic energy is then computed as $k = \omega \nu_t$.

The default, and recommended, values are

$$\omega = 10U_{\infty}/L$$
$$k = \omega \nu_t$$

where the freestream turbulent viscosity ν_t is arbitrarily set to $0.001\nu_l$.

Note that

- If $val_k > 0$, a positive value must be specified for val_om .
- If $val_k \leq 0$, val_om should not be specified.
- If $val_om < 0$, any value specified for val_k is ignored.

For structured grids, inflow turbulence levels may be specified for the SST model in the ARBITRARY INFLOW keyword block. If this is done, the TURBULENCE keyword must come before the ARBITRARY INFLOW keyword block in the input data (.dat) file.

See Also: ARBITRARY INFLOW

Production Term Modification

```
{\tt SST~[TRANSITION]~COEF\_PTM1~} value
```

When TURBULENCE SST TRANSITION is specified, indicating that the compressible version of the SST model with transition modeling is to be used, the separate keyword SST COEF_PTM1 may be used to set the value of the coefficient ϕ_{PTM1} . This coefficient adjusts how much the turbulence production is limited in laminar regions. For a flat plate test case, a value of 1.0 was found to best reproduce the axial location of fully turbulent flow, while a value of 2.0 better predicted the transition onset location. The default value is 1.0.

Chien and Rumsey-Gatski k- ϵ Models³⁷

Several keywords may be used with the Chien and Rumsey-Gatski k- ϵ models to control the initialization procedure, enhance their stability, and improve their accuracy in adverse pressure gradients and at high Mach numbers. For convenience, all keyword phrases associated with the k- ϵ models begin with K-E. Note that all of these apply to structured grids only.

 ${\tt K-E~INITIALIZE~[FROM]~\{\underline{EXISTING}~|~EQUILIBRIUM~|~FREESTREAM\}}$

 $^{^{37}}$ The material in this section was originally written by Dennis Yoder of NASA Glenn Research Center.

This keyword determines how the turbulent transport variables $(k, \epsilon, \text{ and } \mu_t)$ for the k- ϵ models will be initialized.

It is recommended that the user first obtain an intermediate solution using another turbulence model, before switching to one of the k- ϵ models. Initializing from an existing turbulent solution rather than uniform values aids somewhat in convergence and improves the stability of the models by reducing the dramatic changes in turbulence values that occur during the first few iterations after initialization.

This intermediate solution need not be fully converged, but should have reached a state where the the turbulence is well established within the shear regions (boundary layers, mixing layers, etc.) of each zone. Users can examine values of mut/muinf with CFPOST to check the state of turbulence in the flow prior to switching models. Low values of mut/muinf (say < 50 or 100) may be insufficient to sustain turbulence, causing the solution to relaminarize as additional cycles are performed. The SST and Spalart models are good choices for obtaining the intermediate solution.

There are three methods for establishing the initial values for the k- ϵ models.

The first method, given by the EXISTING parameter, performs an "intelligent" initialization, based on the type of turbulence model used in the previous run. This is the default. When initializing from another two-equation model, direct conversion of turbulence values is performed. For lower-order models, the procedure is equivalent to using the EQUILIBRIUM parameter. When initializing from inviscid or laminar solutions, the procedure is equivalent to using the FREESTREAM parameter.

The second method, given by the EQUILIBRIUM parameter, uses an assumption of turbulent equilibrium, namely that the production, Π , of turbulent kinetic energy equals the rate of dissipation, together with an existing turbulent viscosity profile to initialize the k and ϵ variables.

$$\rho \epsilon = \Pi / Re$$

$$\rho k = \sqrt{\frac{\rho \epsilon \mu_t}{C_\mu f_\mu Re}}$$

The third method, given by the FREESTREAM parameter, initializes the turbulence variables to uniform values within each zone.

$$\rho k = \rho k_{\infty}$$
$$\rho \epsilon = \rho \epsilon_{\infty}$$
$$\mu_t = (\mu_t)_{\infty}$$

where the local density is used and the freestream conditions k_{∞} and $(\mu_t)_{\infty}$ may be specified with the keywords K-E FREE_K and K-E FREE_MUT. This method is generally less successful than the others.

These keywords may be used to specify the freestream turbulence values to be used when initializing the turbulence variables to uniform values (i.e., with the K-E INITIALIZE FREESTREAM keyword). The interpretation of the input depends on the signs of val1 and val2.

For the turbulence kinetic energy k_{∞} ,

$$k_{\infty} = \begin{cases} \frac{3}{2}I^{2}u_{\infty}^{2} & \text{for } val1 < 0, \text{ and where } |val1| = I\\ \bar{k}_{\infty}/\bar{a}_{\infty}^{2} & \text{for } val1 > 0, \text{ and where } val1 = \bar{k}_{\infty} \text{ (ft/sec)}^{2} \end{cases}$$

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and for the turbulent viscosity $(\mu_t)_{\infty}$,

$$(\mu_t)_{\infty} = \begin{cases} (\mu_t)_{\infty} & \text{for } val2 < 0, \text{ and where } |val2| = (\mu_t)_{\infty} \\ (\bar{\mu}_t)_{\infty}/\bar{\mu}_{\infty} & \text{for } val2 > 0, \text{ and where } val2 = (\bar{\mu}_t)_{\infty} \text{ (slug/ft-sec)} \end{cases}$$

where the overbar denotes a dimensional value.

If the K-E FREE_K keyword is not used, or if val1=0, then a default value of val1=-0.01 is used, which corresponds to a turbulence intensity I of 1%. If the K-E FREE_MUT keyword is not used, or if val2=0, then a default value of val2=-0.001 is used. This sets the freestream turbulent viscosity to be 0.001 times the freestream laminar viscosity. Note that for values greater than zero, the expected units for val1 and val2 are $(ft/s)^2$ and slug/ft-s, respectively.

These freestream values are only used when the k- ϵ model is initialized from uniform conditions. Currently, the model extrapolates to get values at all inflow and outflow boundaries. Thus, K-E FREE_K and K-E FREE_MUT can not be used to specify inflow conditions.

K-E REINITIALIZE

This keyword signals the code to ignore the old k- ϵ information in the flow (.cfl) file and perform a fresh initialization from uniform values. This command must be removed on subsequent runs or else the model will reinitialize itself each time. Under normal operation, this keyword should not be used.

K-E LINEARIZATION APPROXIMATE | FULL

This keyword controls the implicit treatment of the k- ϵ source terms. The APPROXIMATE formulation results in greater diagonal dominance by neglecting coupling of the k and ϵ equations. The key formulation includes the coupling terms. The default is APPROXIMATE.

K-E [TVD] ORDER
$$\{\underline{1}|2|3\}$$

This keyword sets the spatial order of accuracy of the TVD upwinding used in solving the k- ϵ equations. The default is first order.

```
K-E RELAX [FOR] val [ITERATIONS]
```

Updated values of k, ϵ , and μ_t will be relaxed for val iterations (the default is 500) following the initialization. Relaxation of each of these variables reduces the amount they may change during any single iteration. Immediately after initialization, the allowed changes are significantly reduced. This restriction is then gradually lifted as the last relaxation iteration is approached.

```
K-E [MAXIMUM] [TURBULENT] VISCOSITY val1 K-E [TURBULENT] [REFERENCE] VELOCITY val2
```

The k- ϵ model uses limiters within the interior of each zone to increase convergence and stability by capping the values of the turbulence quantities at both the high and low extremes. This is usually only necessary during the first few iterations after initialization, when the fluctuations in k and ϵ tend to be the most dramatic. Nondimensional values of the minimum limiters have been preset to small numbers. k_{min} is set to 10^{-10} , ϵ_{min} is set to 10^{-12} , and $(\mu_t)_{min}$ is computed from the turbulent viscosity relation using an assumed reference density of 1, $C_{\mu} = 0.09$, and $F_{\mu} = 1$.

The above keywords determine the maximum limiting values for the turbulence quantities. The first keyword sets the maximum turbulent viscosity to be val1 (the default is 10,000) times the freestream viscosity. The second keyword sets the turbulent reference velocity equal to val2 (the default is 1.0) times the freestream speed of sound. The maximum turbulent kinetic energy allowed is 10% of the kinetic energy of the turbulent reference velocity:

$$k_{max} = 0.10 \frac{u_{ref(k-\epsilon)}^2}{2}$$

The maximum dissipation rate is again computed from the turbulent viscosity relation.

The use of these limiters can be summarized as follows:

- If k or ϵ falls below its preset minimum value, it is reset to the minimum value. This typically occurs in the freestream.
- If the turbulent kinetic energy exceeds k_{max} , it is capped at this value. The dissipation rate is taken to be the larger of the current dissipation rate or ϵ_{max} .
- If the turbulent viscosity exceeds $(\mu_t)_{max}$, it is capped at this value and the turbulent kinetic energy is recomputed from the turbulent viscosity relation. The turbulent dissipation rate is left unchanged.

If the maximum limiters cause the turbulence variables to be capped within the flowfield, a warning message will be written to the list output (.lis) file during the final cycle. By using CFPOST to examine the normalized variable mut muinf, one can observe where these limiters are being used and adjust them using the above keywords. It is important that the turbulence values not be limited upon convergence.

K-E COMPRESSIBILITY [CORRECTION] {NONE |
$$\overline{\text{OFF}}$$
 | SARKAR | WILCOX | USER α_k M_{t_0} }

This keyword may be used to specify a compressibility correction, designed to enhance predictions at higher Mach numbers. The equation for the compressibility correction is

$$F(M_t) = \alpha_k \max \left(M_t^2 - M_{t_0}^2, 0 \right)$$

The turbulent Mach number M_t is defined from $M_t^2 = 2k/a^2$, where a is the local speed of sound.

The parameters α_k and M_{t_0} for the various options are defined in the following table.

Correction Type	α_k	M_{t_0}
NONE OFF	0.0	0.00
SARKAR	1.0	0.00
WILCOX	1.5	0.25

Note that both NONE and OFF disable the compressibility correction, and that values of α_k and M_{t_0} should only be included when using the USER option. The default is OFF.

Both the Sarkar and Wilcox compressibility corrections are designed to improve the prediction of compressible jet flows by including the compressible portion of the dissipation rate in the transport equation for the turbulent kinetic energy. These corrections use simple algebraic relations between the solenoidal and compressible dissipation rates. The effect of these corrections is to reduce the

turbulent kinetic energy in regions where the flow is supersonic. In terms of supersonic jet predictions, this results in slower spreading rates, reduced mixing, and a longer core length.

It is well known that the baseline k- ϵ model is poorly suited to adverse pressure gradient flows, such as those found in diffusers. Rodi and Scheuerer (1986) demonstrated that for these types of flows, the rate of dissipation near solid boundaries is too small relative to the rate of production of turbulent kinetic energy. This causes the model to overpredict skin friction and predict flows to be attached when experimental results show them to be separated.

The variable C_{μ} formulation, which is derived from algebraic stress modeling, is designed to help remedy this problem by reducing the turbulent viscosity in regions of the flowfield where the production of turbulent kinetic energy is significantly larger than the rate of dissipation. The specific formulation used is:

$$C_{\mu} = \min\left(0.09, \frac{0.10738(0.64286 + 0.19607R)}{\left[1 + 0.357(R - 1)\right]^2}\right)$$

As the ratio R of production to dissipation increases above 1, the coefficient C_{μ} is reduced from its normal value of 0.09 to limit the turbulent viscosity.

The variable C_{μ} option can provide added stability to the k- ϵ model, such as in the case of an airfoil, where the sudden deceleration of the flow near the leading edge would otherwise result in a significant rate of production. In regions of the flow where the turbulence is in equilibrium, i.e., where the production and dissipation are balanced, the turbulent viscosity remains unchanged.

The above keyword may be used to turn this option on or off (the default is OFF) for the Chien k- ϵ model It has no effect for the Rumsey-Gatski k- ϵ model.

K-E CMUMIN
$$val$$

This keyword controls the lower limit of the C_{μ} coefficient used in the Rumsey-Gatski model. As part of the algebraic Reynolds stress formulation C_{μ} varies throughout the flowfield with typical values in the range [0.07, 0.19]. The default value for this limiter is 0.0005, as recommended by Rumsey.

See Also: TEST 29, TEST 51

Combined RANS/LES Models

The idea behind combined RANS/LES (Reynolds-Averaged Navier Stokes / Large Eddy Simulation) turbulence models is to improve predictions of complex flows in a real-world engineering environment, by allowing the use of LES methods with grids typical of those used with traditional Reynolds Averaged Navier Stokes models. The combined model reduces to the standard RANS model in high mean shear regions (e.g., near viscous walls), where the grid is refined and has a large aspect ratio unsuitable for LES models. As the grid is traversed away from high mean shear regions, it typically becomes coarser and more isotropic, and the combined model smoothly transitions to an LES model.

Several combined RANS/LES models are available in Wind-US. The combined models may only be used for unsteady flows (i.e., the time step is a constant). They are zonal, however, so you can use a combined model in time-accurate mode in one zone, while using a standard RANS model in

steady-state mode in the other zones. For example, a three-zone problem could use the standard SST model with a specified CFL number in zones 1 and 2, and the combined SST/LES model (implemented using an ϵ limiter) with a specified time step in zone 3, using the following keywords:

```
TURBULENCE SST
LESB ZONE 3
CFL# 1.3
TIMESTEP SECONDS 5.0E-6 ZONE 3
```

This capability can accelerate the solution convergence tremendously, especially for large configurations (10 to 20 million grid points) that would be impossible to run in time-accurate mode throughout the flow field.

Spalart DES Model

```
DES [CDES cdes] [zone_selector]
```

When TURBULENCE SPALART is specified, the separate keyword DES may also be input to use the Spalart Detached Eddy Simulation (DES) turbulence model. It is intended to improve the results for unsteady and massively separated flows. The DES model reduces to the standard Spalart-Allmaras model near viscous walls, where the grid is refined and has a large aspect ratio, but acts like a Large Eddy Simulation (LES) model away from the boundary, where the grid is coarser and has an aspect ratio of order one.

The input parameter cdes specifies the value of the coefficient C_{des} in the model. The default value is 0.65. Increasing C_{des} increases the size of the region in which the DES model reduces to the standard Spalart-Allmaras model. Details may be found in papers by Spalart, Jou, Strelets, and Allmaras (1997) and by Shur, Spalart, Strelets, and Travin (1999).

Shih's Modified DES Model

```
MDES [CDES cdes] [zone_selector]
```

The MDES keyword is similar to the DES keyword described above, and may be specified in conjunction with TURBULENCE SPALART to use Shih's modified version of the DES model. As in the standard DES model, the input parameter cdes specifies the value of the coefficient C_{des} , and the default value is 0.65.

SST with ϵ Limiter

```
LESB [CBLES cb] [zone\_selector]
```

The LESB keyword may be used with the SST model (TURBULENCE SST) to specify use of a combined SST and LES turbulence model, using an ϵ limiter.

The input parameter cb specifies the value of the coefficient C_B in the model. The default value is 10.0. Increasing C_B increases the size of the region in which the combined model reduces to the standard SST model.

See Also: TEST 15

Nichols-Nelson Hybrid Model

$\verb|HYBRID [VERSION ver] [CLES $cles$] [DELTA $grid_scale$] [zone_selector]|\\$

The HYBRID keyword may be used with either the SST or the Chien k- ϵ model to specify use of the Nichols-Nelson hybrid RANS/LES turbulence model (Nichols and Nelson, 2003).

ver A flag indicating the version of the model to be used.

- 1 The original Nichols-Nelson hybrid model. The μ_t value from the RANS model is used in the turbulence model equations, and the hybrid μ_t value is used in the Navier-Stokes equations.
- 2 The hybrid μ_t value is used in both the turbulence model and Navier-Stokes equations.
- 3 The hybrid μ_t value is used in both the turbulence model and Navier-Stokes equations, but k is treated like a sub-grid kinetic energy, not the full turbulent kinetic energy. (I.e., the model becomes more of an LES subgrid model.)

The default value is 1.

cles The coefficient c_{LES} used when calculating the LES value of the turbulent viscosity.

$$(\mu_t)_{\rm LES} = c_{\rm LES} \, \rho \, \delta \, k^{1/2}$$

The default value is 0.0854.

grid_scale A flag indicating the version of the model to be used.

1 $\delta = \max(dx, dy, dz)$

2 $\delta = \text{volume}^{1/3}$

The default value is 1.

Partially Resolved Numerical Simulation (PRNS)

PRNS [RCP rcp] [zone_selector]

The PRNS keyword may be used with the Spalart-Allmaras and SST models to specify use of the PRNS (Partially Resolved Numerical Simulation) model, and is only applicable to unstructured grids. The input value Rcp is the resolution control parameter, the ratio of the temporal filter width to the turbulent integral time scale. A value of Rcp = 1.0 corresponds to a Reynolds-averaged solution (i.e., no turbulent eddy resolution), while lower values result in smaller turbulent eddies being resolved. The default value is 0.4, intended to correspond to a very large eddy simulation.

Note that lowering Rcp means that more grid points are required to resolve the smaller eddies. In their original paper describing the PRNS model, Liu and Shih suggest that the number of grid points for a PRNS solution N_{PRNS} may be estimated using the formula

$$N_{\rm PRNS} = N_{\rm RANS} Rcp^{-9/4}$$

where N_{RANS} is the number of grid points required for a Reynolds-averaged solution.

For more details on the PRNS model see the papers by Liu and Shih (2006) and by Shih, Liu, and Chen (2006).

Detached PRNS

 ${\tt DETACHED-PRNS} \ \ [{\tt RCP} \ \ rcp] \ \ [{\tt DPRNS} \ \ dprns] \ \ [{\tt zone_selector}]$

This keyword is similar to the PRNS keyword described above, but may only be used with the Spalart-Allmaras model. It specifies use of the Detached PRNS model, and is also only applicable to unstructured grids. The input value Rcp is again the resolution control parameter, with a default value of 0.4.

Unlike the standard PRNS model, the Detached PRNS model doesn't scale the production term and turbulent viscosity near viscous walls. The size of this "near-wall" region is given by

$$D_{PRNS}V_{cell}^{1/3}$$

where D_{PRNS} is the user-specified value dprns, and V_{cell} is the local cell volume. The default value for dprns is 5.0.

TVD — Total Variation Diminishing operator flag

Structured Grids

```
TVD [OFF | FACTOR factor | MINMOD [factor] | KOREN [factor] | ALBADA] \
    [zone_selector]
```

Unstructured Grids

```
TVD [OFF | FACTOR factor | MINMOD [factor] | SUPERBEE factor | \
    VENKAT [factor [KVENKAT K_{Venk}]]] [zone\_selector]
```

Structured Grids

For structured grids, this keyword controls the TVD flux limiter in the explicit operator, which is helpful in preventing overshoots in flowfield properties in regions of high gradients. TVD may be used with the Coakley, Roe, Van Leer, HLLC, HLLE, and Rusanov explicit schemes. TVD is enabled by default for these schemes, using the standard minmod TVD limiter with the default compression parameter, and disabled for all others.

OFF Disables TVD for the specified zone(s)

Use the standard minmod TVD limiter. The value of factor specifies the "com-FACTOR pression" parameter for the TVD algorithm, which controls the amount of limiting applied.

Use the standard minmod TVD limiter. This is the same as FACTOR, except that the MINMOD factor may be defaulted. The value of factor specifies the "compression" parameter for the TVD algorithm, which controls the amount of limiting applied. The default compression parameter is the maximum value which assures TVD properties for the explicit operator being used (as specified by the RHS keyword). The default values are shown in the following table for the various versions of the explicit differencing scheme.

Explicit Operator	factor
First-order, upwind	0 (i.e., not applicable)
First-order, upwind, modified for stretched grids	2
Second-order fully upwind	2
Second-order central	2
Second-order upwind-biased	3
Second-order upwind-biased, modified for stretched grids	3
Third-order upwind-biased	4
Third-order fully upwind	Not available
Fourth- and fifth-order	Not available

KOREN Use the Koren TVD limiter. With this limiter, the default value of factor is 1.0, and does not depend on the explicit operator being used.

ALBADA Use the van Albada TVD limiter. With this limiter, no factor value is input.

If TVD flux limiting is turned on for an invalid explicit scheme, but without specifying a compression parameter, TVD flux limiting will automatically be disabled and the run will continue. Otherwise, an error message is generated and the run will abort.

See Also: BOUNDARY TVD, RHS

Unstructured Grids

For unstructured grids, standard slope limiting is performed by default for all viscous cases, and for inviscid cases with second-order spatial differencing specified with the RHS keyword.

OFF Disables slope limiting for the specified zone(s)

FACTOR Use the standard slope limiting procedure. The value of factor specifies the slope

limiting parameter, and must be specified.

MINMOD Use the standard slope limiting procedure. This is the same as FACTOR, except

that the factor may be defaulted. The default slope limiting parameter is 0.0 for

first-order spatial differencing, and 3.0 for second-order differencing.

SUPERBEE Use "super-bee" slope limiting. The slope limiting parameter factor must be speci-

fied.

VENKAT Use Venkatakrishnan slope limiting (Venkatakrishnan, 1995). A value of 3.0 for the

slope limiting parameter *factor* corresponds to second-order, and 0.0 corresponds to first-order. Values between 0.0 and 3.0 result in linear interpolation between the first-order and second-order face states. The default (and maximum allowed) value

for factor is 3.0.

The optional K_{Venk} specified using KVENKAT is a coefficient determining the value of a "knob" that controls the allowable amount of oscillation. The value is set to $K_{Venk}D_{cell}$, where D_{cell} is the diameter of a sphere with the same volume as the

current cell. The default for K_{Venk} is 20.

See Also: CONVERGE LOAD, RHS

VISCOSITY — Specification of viscosity law

Structured Grids

 $\mbox{VISCOSITY } \{ \mbox{\underline{SUTHERLAND}} \ | \ \mbox{WILKE} \ | \ \mbox{KEYE} \ | \ \mbox{CONSTANT} \ vis \ | \ \mbox{TUNNEL9} \ | \ \mbox{CUSTOM} \ c_1 \ c_2 \}$

Unstructured Grids

VISCOSITY {<u>SUTHERLAND</u> | WILKE}

This keyword allows you to specify the method of computing the transport properties.

The equations shown below are for the laminar viscosity coefficient μ . For all the options except WILKE, in Wind-US the laminar thermal conductivity coefficient k is equal to the viscosity coefficient, when non-dimensionalized. For WILKE, the form of the equations used for k is the same as those used for μ , but with different constants from the chemistry data (.chm) file.

In all of the equations, μ is in slug/ft-sec and T is in ${}^{\circ}\mathbf{R}$.

Structured Grids

SUTHERLAND Use Sutherland's law, designed for ideal gases with T > 180 °R, as follows:

$$\mu = 2.329 \times 10^{-8} \frac{T^{3/2}}{T + 216}$$

This is the default.

WILKE Use Wilke's law, designed for multi-species flow (real gases). First, the viscosity coefficient is computed for each individual species n using Sutherland's law, as follows:

$$\frac{\mu_n}{\mu_0} = \left(\frac{T}{T_0}\right)^{3/2} \frac{T_0 + S}{T + S}$$

where T is the local static temperature, and μ_0 , T_0 , and S are constants read from the chemistry data (.chm) file for species n. For N total species, the individual viscosity coefficients are combined using

$$\mu = \sum_{i=1}^{N} \frac{X_{i} \mu_{i}}{\sum_{j=1}^{N} X_{j} \phi_{i,j}}$$

where $\phi_{i,j}$ is a mixing coefficient computed as

$$\phi_{i,j} = \left[8 \left(1 + \frac{M_i}{M_j} \right) \right]^{-1/2} \left[1 + \left(\frac{\mu_i}{\mu_j} \right)^{1/2} \left(\frac{M_j}{M_i} \right)^{1/4} \right]^2$$

X is the species mole fraction, and M is the species molecular weight.

KEYE

Use Sutherland's law for $T \ge 180$ °R, Keyes' law for $T \le 160$ °R, and a linear combination for 160 °R < T < 180 °R. Sutherland's law is written as above. Keyes' law is given by:

$$\mu = 2.32 \times 10^{-8} \frac{T^{1/2}}{1 + (220/T) \times 10^{-9/T}}$$

And the linear combination is given by

$$\mu = f\mu_S + (1 - f)\mu_K$$

where μ_S and μ_K are the viscosity coefficients from Sutherland's and Keyes' laws, and f = (T - 160)/20.

CONSTANT vis Use a constant molecular viscosity of vis (slug/ft-sec)

TUNNEL9 Use a viscosity model obtained from AEDC Tunnel 9.

$$\mu = \begin{cases} 8.913 \times 10^{-10} \, T^{\, 0.979} & \text{for } T \leq 227 \, ^{\circ} \mathrm{R} \\ 2.144 \times 10^{-8} \, T^{\, 3/2} / (T + 179.1) & \text{for } 227 \, ^{\circ} \mathrm{R} < T \leq 795 \, ^{\circ} \mathrm{R} \\ 6.050 \times 10^{-9} \, T^{\, 0.659} & \text{for } T > 795 \, ^{\circ} \mathrm{R} \end{cases}$$

CUSTOM c_1 c_2 Use Sutherland's law with the constants c_1 and c_2 , as follows:

$$\mu = c_1 \frac{T^{3/2}}{T + c_2}$$

Unstructured Grids

For unstructured grids, only Sutherland's and Wilke's laws are available. The equations are the same as shown above for structured grids.

VORTEX GENERATOR — Vortex generator model (block)

Structured Grids

```
VORTEX [GENERATOR]

{MODEL WENDT

ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \

[SUBSET I range J range K range]

ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \

[SUBSET I range J range K range]

NUMBER ival

vg_boundary {XLOC xl | YLOC yl | ZLOC zl} chord height alpha \

[VEL vmax] [DEL delta] |

MODEL BAY

ZONE izone

NUMBER ival

VANE_SPEC imin imax jmin jmax kmin kmax vg_boundary alpha area \

[model_constant]}

ENDVORTEX
```

This keyword block is applicable to three-dimensional cases with structured grids, and enables the user to specify the effects of vortex generators in the flow using a choice of two models: the Wendt model (Dudek, 2006; Wendt, 2001) and the BAY model (Bender, Anderson, and Yagle, 1999). Both models simulate the vortices produced by an array of vane-type vortex generators. A vortex generator array consists of one or more vortex generators mounted on viscous wall boundaries. A separate vortex generator keyword block must be used for each array. The models are only applicable to generators placed on viscous wall boundaries. It is the user's responsibility to check this — Wind-US currently has no mechanism for flagging this input error.

The various elements of the VORTEX GENERATOR input block are defined as follows:

```
VORTEX [GENERATOR]
```

Defines the beginning of the vortex generator keyword input block.

```
ENDVORTEX
```

Ends the vortex generator input block

```
{MODEL WENDT | MODEL BAY}
```

The main part of the VORTEX GENERATOR block is divided into two sections, corresponding to input for the Wendt and BAY models. Either MODEL WENDT or MODEL BAY must be specified, but not both.

Wendt Model

The Wendt model uses a discontinuous change in secondary velocity across a zonal interface boundary in order to simulate the vortices produced by an array of vane-type vortex generators. The model determines the strength of each vortex based on the generator chord length, height and angle of incidence with the incoming flow, as well as the incoming flow core velocity and boundary layer thickness. Each vortex center is placed at the grid point closest to the location determined by the user-specified generator location on the boundary, and the generator height.

Note that the Wendt vortex generator model is derived from experimental data taken at axial stations which were one chord length downstream of the trailing edge of the generators, and therefore the secondary velocities produced by the model simulate vortices at this station, rather than at the generator trailing edges. For most accurate results, the boundary where the vortex generator boundary condition is applied should be at this one chord length station.

The following restriction applies:

• The BOUNDARY TVD FACTOR 0 keyword option should be used at all interface boundaries containing vortex generators.

The elements of the VORTEX GENERATOR input block applicable to the Wendt model are defined as follows:

```
ZONE iz1 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \ [SUBSET I range J range K range]
ZONE iz2 BOUNDARY {I1 | IMAX | J1 | JMAX | K1 | KMAX} \ [SUBSET I range J range K range]
```

These two lines define the location of the vortex generator array. The relevant zones are given by the values of iz1 and iz2, and the relevant boundaries within zones iz1 and iz2 are specified via the BOUNDARY keyword parameter.

- iz1 The "upstream" zone. The secondary velocities will be increased when passing information from this zone to iz2.
- iz2 The "downstream" zone, i.e., the zone receiving increased secondary velocities from zone iz1. Secondary velocities will be decreased when passing information from this zone back to zone iz1.

Note: Currently, the vortex generators must be located at either an i_1 or i_{max} boundary. Thus, the only valid choices with the BOUNDARY keyword parameter are I1 and IMAX.

Note that the secondary velocity is the flow in a plane normal to the primary velocity. For example, for a vortex generator array at an i-interface boundary, the primary flow is in the i direction, and the secondary velocity is in the j and k directions.

The SUBSET parameter may be used to specify that the change in secondary velocity occurs only over a part of the zone boundary. Otherwise, it is assumed that the change occurs over the entire boundary. The *range* parameters define the part of the zone boundary over which the change occurs, and take one of the following forms:

index1 index2 Starting and ending indices in the specified direction. LAST may be used for the last index.

ALL Equivalent to 1 LAST.

The starting and ending indices for the appropriate I, J, or K parameter (depending on the boundary specified) must be the same, and correspond to that boundary.

NUMBER ival

This defines the number of vortex generators in the vortex generator array (i.e., on the specified boundary).

```
vg\_boundary {XLOC xl | YLOC yl | ZLOC zl} chord\ height\ alpha \
   [VEL vmax] [DEL delta]
```

Defines the location and geometric parameters of each vortex generator. A separate line of input must be included for each generator.

 $vq_boundary$

The minimum or maximum index, specified as IM, IX, JM, JX, KM, or KX, within the specified boundary surface (as specified by ZONE and SUBSET) where the generator is mounted. This boundary must have been defined as a viscous wall. As an example, for a vortex generator array at an iinterface boundary, either JM, JX, KM, or KX should be specified, where JM means the generator is at the j=1 interface boundary, JX means it's at the $j = j_{max}$ boundary, etc.

XLOC, YLOC, ZLOC

xl, yl, or zl is the x, y, or z coordinate location in inches of the base of the vortex generator on the already specified boundary surface and wall boundary. (Currently, only one coordinate direction and location may be specified. This may lead to ambiguity in specifying the locations for generators placed in complex duct geometries. This shortcoming will be addressed in future code updates.)

chord

The chord length of the vortex generator in inches. Must be greater than zero.

height

The height of the vortex generator in inches. Must be greater than zero.

alpha

The angle of incidence of the generator in degrees. Must be non-zero. More specifically, it is defined as the angle the generator chord line makes with the primary flow direction. The sign of alpha is determined using the "right-hand rule" of vector mechanics. For a given generator, use a normal vector pointing into the wall (the thumb), and the primary flow direction (fingers). If the rotation from the primary flow direction towards the generator follows the right-hand rule, then the sign of alpha is positive. Otherwise, it is negative. A positive value of alpha generally results in a vortex with a counterclockwise rotation, and conversely. The recommended range of magnitudes for alpha is between 8 and 20°.

VEL

vmax is the maximum velocity at the vortex generator station, in ft/sec. The default is for Wind-US to compute this value from the flow. It is currently recommended that this value be specified by the user. Must be

greater than zero.

DEL

delta is the boundary layer thickness of the incoming flow, in inches. The default is for Wind-US to compute this value from the flow. It is currently recommended that this value be specified by the user. Must be greater than zero.

In selecting values for the above described parameters, keep in mind that the vortex generator model was derived from conservation of momentum and inviscid theory, correlated with experimental data having height/chord ratios between 0.13 and 2.62, height/delta ratios between 0.12 and 2.60, and for duct flows with core Mach numbers ranging from 0.20 to 0.60. Because of the theory used in its derivation, the model is intended to work well outside of the range of the experiments. However, there is one caveat: past experience indicates that the model does not work when the vortex generators are placed in regions of sonic flow.

Example

The following example illustrates the use of the Wendt vortex generator model for one vortex generator array located between zones 2 and 3. The interface between zones 2 and 3 corresponds to the i_{max} boundary of zone 2 and the i_1 boundary of zone 3. The array contains two vortex generators, both mounted on the j_{max} viscous wall boundary of the zone interface with z coordinate values of -0.52 inches and 0.52 inches. The generators have the same geometric parameters: the chord length is 1.6 inches, the height is 0.4 inches, and the angle of incidence is 16° .

```
VORTEX GENERATOR
MODEL WENDT
ZONE 2 BOUNDARY IMAX
ZONE 3 BOUNDARY I1
NUMBER 2
JX ZLOC -0.52 1.6 0.4 16.0
JX ZLOC 0.52 1.6 0.4 16.0
ENDVORTEX
```

See Also: BOUNDARY TVD

BAY Model

The BAY model is a source term model which models the side force produced by the vortex generator and adds it to the momentum and energy equations. This side force automatically adjusts its strength based on the local flow. The user specifies the grid points over which the force is to be applied (i.e., enclosing each vortex generator).

The elements of the VORTEX GENERATOR input block applicable to the BAY model are defined as follows:

```
ZONE izone
```

This line defines the zone containing the vortex generator array.

```
{\tt NUMBER}\ ival
```

This defines the number of vortex generators in the vortex generator array.

```
{\tt VANE\_SPEC} \ imin \ imax \ jmin \ jmax \ kmin \ kmax \ vg\_boundary \ alpha \ area \ [model\_constant]
```

This keyword line must be specified *ival* times, i.e. once for each vane. It is used to specify the grid points over which the model acts for each vane and the accompanying vane angle of incidence, planform area and model constant. These parameters are described in more detail below.

imin, imax Minimum and maximum i indices defining the grid region enclosing the vortex generator

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jmin, *jmax* Minimum and maximum *j* indices defining the grid region enclosing the vortex

generator

kmin, kmax Minimum and maximum k indices defining the grid region enclosing the vortex

generator

vq_boundary The minimum or maximum index, specified as IM, IX, JM, JX, KM, or KX, indi-

cating the viscous wall where the generator is mounted. Here IM means the vane is on the i=1 boundary, IX means it's on the $i=i_{max}$ boundary, etc.

alpha The angle of incidence of the vane chord line (in degrees) with respect to the

coordinate grid line in the primary flow direction. Note: In instances where the leading edge of the vane corresponds to the maximum index of the VANE_SPEC range rather than the minimum (i.e., jmax or kmax approximately specify the leading edge of the vane), the sign of the angle must be negative in order to

produce the correct directional rotation of the vortex.

area The vane area (in^2)

model_constant The model constant. The default value is 10.

Example

The following example illustrates the use of the BAY vortex generator model for one vortex generator array located in zone 2. The array contains two vortex generators, both mounted on the j_{max} viscous wall boundary. The VANE_SPEC values of the zones are shown below. They both act over the same i and j indices but at different k indices on the wall. Both generators have the same angle of incidence with the primary flow direction grid line, 16° , and have planform areas of 0.21 in².

```
VORTEX GENERATOR

MODEL BAY

ZONE 2

NUMBER 2

VANE_SPEC 15 25 45 65 20 25 JX 16.0 0.21
```

VANE_SPEC 15 25 45 65 20 25 JX 16.0 0.21 VANE_SPEC 15 25 45 65 35 40 JX 16.0 0.21

ENDVORTEX

WALL FUNCTION — Specify the use of wall functions

 $\verb|WALL FUNCTION {ON|OFF}| [zone_selector]|\\$

For turbulent flows, this keyword may be used to invoke wall function boundary conditions on viscous walls, using the White-Christoph law of the wall. Wall function boundary conditions allow calculations to be performed with fewer grid points, and generally with higher CFL numbers.

A wall function boundary must lie on one of the boundaries of the selected zone, and may be an overlapped or internal boundary. The first grid point adjacent to the boundary must be within the log layer (roughly $15 < y^+ < 100$). The recommended y^+ value at the first point off the wall is about 50. The boundary condition will revert to a no-slip condition when the first grid point off the wall falls below $y^+ = 15$, indicating that it is within the laminar sublayer.

Note that the wall function boundary condition is used to eliminate the grid points that would otherwise be required to resolve the laminar sublayer and the logarithmic layer. The grid beyond the first point off the wall should be the same as for a run without wall functions, in order to properly resolve the rest of the boundary layer. For structured grids, the *cfsubset* utility may be useful when modifying an existing grid for use with wall functions.

The implementation of wall functions in Wind-US involves only the modification of the wall flux, and does not reset the values of any flow quantities.

Viscous forces on wall function boundaries calculated using the LOADS keyword should only be used for determining convergence. For structured grids, correct values can be calculated from the .cfl file using the integrate force command in CFPOST.

Based on limited test cases, the use of wall functions is not recommended if accurate predictions of wall heat transfer are required, or for flows with strong shock waves.

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WALL SLIP — Iterations until no slip

Structured Grids

```
WALL SLIP [ITERATIONS] val [zone_selector]
```

To minimize transients at the start of a Wind-US calculation, with structured grids the velocity at no-slip boundaries is actually reduced from its initial value to the no-slip condition over a number of iterations. The WALL SLIP keyword allows the user to specify the number of iterations, given by val, on a zonal basis. The default value is 50 iterations.

WALL TEMPERATURE — Specify wall temperature

WALL TEMPERATURE { $\underline{ADIABATIC} \mid value \mid \texttt{EQUILIBRIUM} \mid \texttt{EMISSIVITY} \mid emiss$ } \ [zone_selector]

This keyword allows you to specify a wall temperature on a zonal basis.

ADIABATIC Use an adiabatic wall boundary condition for temperature. This is the default if

the WALL TEMPERATURE keyword isn't used.

value Use a constant wall temperature equal to value °R. Zero-gradient extrapolation is

used to get the chemical species at the wall for finite-rate chemistry.

EQUILIBRIUM [EMISSIVITY] emiss

Set the wall temperature based on thermal equilibrium (i.e., conduction to and

radiation from the wall are equal), with the wall emissivity given by emiss.

See Also: TTSPEC

WRITE — Write extra variables into .cfl file

WRITE [VARIABLES] var [zone_selector]

Using this keyword, additional variables may be computed and written into the .cfl file. The user-specified input var must be one of the keywords listed below. Multiple WRITE keywords may be used to write more than one extra variable into the file.

Skin friction coefficients in the x, y, and z directions. In the .cfl file, the variable names are cfx, cfy, and cfz. Only available for unstructured grids.

LESWIDTH LES indicator for the combined RANS/LES turbulence models; 1 if in LES region, 0 otherwise. This option applies to all the combined RANS/LES models except the Nichols-Nelson hybrid model. In the .cfl file, the variable name is LESwidth. For backward compatibility, this keyword may also be specified as FILTER.

GRADQUAL Gradient quality measures. In the *.cfl* file, the variable names are GradQual1 through GradQual5. Only available for unstructured grids.

HEATFLUX Heat transfer coefficient. In the .cfl file, the variable name is HeatFlux.

RESIDUAL Individual equation residuals. In the .cfl file, the variable names are Resid_rho, Resid_rho*u, Resid_rho*u, Resid_rho*w, and Resid_rho*e0. Only available for unstructured grids.

TOTALS Total pressure and temperature. In the .cfl file, the variable names are Pt-frozen and Tt-frozen.

VENKATPHI Limiter used for Venkatakrishnan slope limiting. In the .cfl file, the variable name is VenkatPhi. Only available for unstructured grids.

11 Test Options

Several user-controlled options have been provided as an aid to modifying Wind-US. These options all may be selected by using the keyword TEST in the input data file. The test options typically control program features which are under test, and have not been accepted for production use. Each test option is of the form

 ${\tt TEST}$ number mode

If mode is not described for a given test option, the user should use mode = 1 to activate that option.

As test options are accepted, they are "hard wired" into the code and the test option described here becomes meaningless. Test options may also be rejected based on trial runs. In that case, the option code described here will also become meaningless. For this reason, the list of valid options is not consecutive. Selection of an invalid option will be accepted by Wind-US, but will have no effect.

The various test options and modes are described in Table 15. For each option, the subroutines and functions referencing that option are listed in parentheses.

Table 15: Non-Production Test Options

number	Description
2	Designed for parallel processing data transfer debugging.
	$\underline{mode} \underline{\text{Result}}$
	1 Don't read zonal boundary data (evrwbd)
	2 Not used
	4 Don't solve zone (evsolv)
	8 Don't write zonal data (evwzon)
	16 Don't write zonal boundary data (evrwbd)
	32 Not used
	64 Not used
	128 Not used
	256 Don't update boundary conditions (structured grids only) (lpschm)
	Set <i>mode</i> equal to the sum of the desired actions.
3	Parallel processing task tracing.
	$mode { m Action}$
	1 Trace event reads/writes (rwev)
	2 Trace file I/O (rwnh, rwc, rwd, rwi, rwr)
	4 Trace network traffic (psexit, psgscat, psrwev, psrwgv, psrwnh,
	${ t psrwc}$, ${ t psrwd}$, ${ t psrwr}$, ${ t rwbc}$
	8 Trace task begin/end (psspwn, tskbeg, tskidl)
	16 Print task queue for debugging (psqprt)
	Set $mode$ equal to the sum of the desired actions. I.e., setting $mode = 5$ will trace both event reads/writes and network traffic.

 Table 15: Non-Production Test Options (Continued)

number	Description
5	Flag for procedure used in gas1 to compute effective specific heat ratio β and sound speed a for frozen and finite-rate chemistry.
	\underline{mode} Procedure
	> 0 Iterate on pressure, with a maximum of <i>mode</i> iterations < 0 Iterate on temperature, with a maximum of 20 iterations
	The default is to iterate on pressure, with a maximum of 20 iterations. (gas1)
6	Write $.cfl$ file compatible with the "old" code. (asnsx, asvisc)
7	Do not use high performance C I/O interface. (openf)
8 9	Use Version 2 common files. (cftopn, mpinit, openf, zhinit) For $mode = 1$, run a verification case using the method of manufactured solutions. If $mode = 2$, in addition to running the case, Plot3d function files named $mms.exact.fxn$, $mms.cfd.fxn$, and $mms.error.fxn$ are written containing the exact solution, the Wind-US solution, and the error.
	The functional forms for the primitive variables are combinations of sine and cosine functions, and are coded in subroutines mms_3d_euler_bc and mms_3d_ns_bc for the Euler and Navier-Stokes equations, respectively. The corresponding source terms for the governing equations are coded in subroutines mms_3d_euler_src and mms_3d_ns_src.
	The choice of an Euler or Navier-Stokes solution is determined by the TURBULENCE keyword. If the flow is inviscid in all zones, an Euler solution is computed; otherwise, a constant-viscosity laminar Navier-Stokes solution is computed. The flow may be 2-D or 3-D, as determined by the grid file.
	Multi-zone grids may be used, but the use of COUPLING MODE CHARACTERISTIC is recommended for Navier-Stokes solutions. (Viscous terms are neglected in Roe boundary coupling, and they are <i>large</i> in the method of manufactured solutions.)
	Frozen boundary conditions must be specified in the grid file at all non-coupled boundaries. The appropriate boundary conditions for the functional forms being used will then be set automatically.
	See Roach (1998) and Salari and Knupp (2000) for more detailed information on the method of manufactured solutions. (12norm1d, lpschm, mms_test_soln, nsrhss, pstinp, sins, tdbcni)
10	Print the time step information (i.e., minimum/maximum CFL and Δt) into the <i>.lis</i> file every <i>mode</i> cycles, instead of just on the first cycle. (NSzsolv)
11	Normally, the flowfield is rotated to be consistent with changes in the global angles of attack and yaw. This TEST option prevents that rotation. (Changes in the global angles may occur at the start of a restart run, if the user specifies angles that are different from the values in the .cfl file, or during a run when the FIXED_CL keyword is used.) (lpgrp)
12	In the HLLC scheme, use a more accurate method for computing the wave speeds. (hllc, US_HLLC, US_HLLCRL)

 Table 15: Non-Production Test Options (Continued)

number	Description
13	Apply a constant grid acceleration of $1000 \times mode $ ft/sec ² in the $+x$ direction. If $mode < 0$, the grid velocity components will be written to the $.cfl$ file, with variable names ui, vi, and wi. Only applies to structured grids, with a time step in seconds or Newton iteration (i.e., without local time steps). May not be used with any other moving grid option. (lprk, pstinp, updatg)
14	For structured grids, add pressure to the .cfl file in read-only mode. Otherwise it's added in read-write mode. (It's always added in read-write mode for unstructured grids.) (axflow)
15	For the SST model with an ϵ limiter (the LESB keyword), modify the grid filter width Δ used in the limiter. This test option only applies to structured grids. (sst1, sst2, sst3)
	$ \frac{mode}{0} \frac{\text{Grid Filter Width}}{\max(ds_1, ds_2, ds_3, V dt, \sqrt{k} dt)} \\ 1 \max(ds_1, ds_2, ds_3, \sqrt{k} dt) \\ 2 \max(ds_1, ds_2, ds_3, V dt) \\ 3 \max(ds_1, ds_2, ds_3) $
	where ds_1 , ds_2 , and ds_3 are the distances along the grid lines in the three directions, V is the velocity, k is the turbulent kinetic energy, and dt is the time step size.
16	Manually set the blowing Mach number for BLOW SURFACE boundary conditions to $mode/100$. The default is to compute the blowing Mach number from the local flow conditions and the specified plenum conditions, with a maximum value of 1.0. (Note that this sets the actual blowing Mach number, not a different maximum value.) (BC_bcbled)
17	Use "new" Baldwin-Barth turbulence model. (bbarth)
20	Non-dimensionalize k and ω in the SST model the "old" way. (aijkrg, aikeps, sstprtinp, sstpstprt)

 Table 15: Non-Production Test Options (Continued)

number	Description
21	Flag affecting the initialization and freestream value of the eddy viscosity for the Spalart-Allmaras and Goldberg turbulence models. (goldbergprtinp, goldbergpstprt, redimsa, sabound, saprtinp, sapstprt, sinut, spalart, TURB_goldbergboundUS, TURB_saboundUS)
	\underline{mode} \underline{Result} $\underline{0, 1}$ Use a default freestream value for $(\nu_t)_{\infty}$ of 5.0, and initialize the dependent variable to $(\nu_t)_{\infty}/\nu_l$. 2 Use a default freestream value for $(\nu_t)_{\infty}$ of 0.1, and initialize the dependent variable to $(\nu_t)_{\infty}$.
	In addition, for the Spalart-Allmaras model in structured grids,
	$\frac{mode}{0}$ Result Use the original 1992 model, with an f_{t2} term for laminar stabilization. Like mode 0, except with "corrections" to the production and destruction terms. This is equivalent to the default model for WIND beta versions 4.15 to 4.92. (Note that's "WIND," which predates "Wind-US".)
	Like mode 1, except without the f_{t2} term. This is equivalent to the default model for WIND versions prior to WIND beta 4.15, and includes a slight error that makes the model overly dissipative.
25	In the Baldwin-Lomax model, use y^+ based on wall vorticity. (blomax)
26 29	Use local values in y^+ damping for the Baldwin-Lomax, Cebeci-Smith, Baldwin-Barth, and k - ϵ models. (bbdamp, blomax, cebeci, kepy2) For the Cebeci-Smith model, use y^+ based on wall vorticity. (algtur, cebeci)
	For the k - ϵ models, TEST 29 is a production limiter (kelhssch, kelhssrg, keprod, kerhssch, kerhssrg, redimkeps3d)
	$mode$ Result0Production limited to $20 \times$ dissipation1Production computed from vorticity, but not limited2Production not limited3Production computed from vorticity, then limited4Production computed using $2\mu_t S_{ij} S_{ij}$, but not limited5Production computed using $2\mu_t S_{ij} S_{ij}$, then limited
	Modes 4 and 5 only apply to the Rumsey-Gatski ASM $k\text{-}\epsilon$ model.
30	Irrotational boundary condition at freestream inflow boundaries (structured zones only). (BC_bcfree)
31	Use quadratic curve fit when applying normal derivative boundary conditions $(\partial p/\partial n \text{ and } \partial T/\partial n)$ at walls. (BC_bcwall)
37	In wbnd2, deallocate memory used for boundary coupling data before returning. (wbnd2)

Table 15: Non-Production Test Options (Continued)

number	Description
40 46	No longer valid; use K-E COMPRESSIBILITY CORRECTION instead. (keppstin) In the SST turbulence model, in blowing regions and bleed regions with a specified negative bleed flow rate, set $\mu_{turb} = 10 \mu_{lam}$ along the wall. (sstbound, TURB_sstboundUS)
47	For the algebraic turbulence models, smooth turbulent viscosity in each <i>i</i> -plane using simple averaging, with the number of smoothing passes set to <i>mode</i> . (smtvis)
48	For the algebraic turbulence models, smooth turbulent viscosity in three dimensions using simple averaging, with the number of smoothing passes set to mode. (smtvis)
49	Modified smoothing for explicit operator for structured grids (see the SMOOTHING keyword). (bdload, dampi, dampj, dampk)
	 mode Result No pressure switch on second-order dissipation No pressure switch, and an LES type filtering of nonlinear terms
51	Limit the turbulent viscosity μ_T such that the maximum value of $\mu_T/(\mu_L)_{\infty} = mode \times 1000$. Suggested range is $50 < mode < 100$. Not valid with the k - ϵ models; use K-E MAXIMUM TURBULENT VISCOSITY instead. (keppstin, mutlim, TURB_mutlim, TURB_UpdateGoldberg, TURB_UpdateSA, TURB_UpdateSST)
52	When using BLOW PLENUM, print a warning when the plenum total pressure is automatically raised because it was less than the local static pressure. (BC_bcbled)
54	Reserved for use at Boeing.
55	Reserved for use at Boeing.
56	No energy addition to fluid due to MFD equations. (emdef)
57	When an explicit solution procedure is being used for the mean flow equations, still solve the Spalart-Allmaras and SST turbulence model equations using an implicit method. This test option only applies to structured grids. (spalart, sst1, sst2, sst3)
58	For MFD flows, read the Lorentz force directly from the .cfl file, but using the .cfl file variable names normally used for the electric field. Requires EFIELD CFL in the MFD keyword block. (emdef)
59	Apply SMOOTHING keyword values to the Spalart-Allmaras model as well as the mean flow solver. (Currently deactivated.) (spalart)
60	Print warning when local and coupled normal velocities have inconsistent directions at a coupled boundary. (postrbs)
61	When $mode=2$, all boundary conditions are applied, whether or not they're consistent with the IBLANK values. This only affects corners, where there are usually multiple boundary conditions. So, if a wall boundary at $j=1$ meets an outflow boundary at $i=i_{max}$, if TEST 61 2 is specified, both boundary conditions are applied. (tdbcgs)

 Table 15: Non-Production Test Options (Continued)

number	Description	
62	Compute cell areas and volumes using procedure from WIND 4. This test option only applies to structured grids. (mphzmet, nsrhsv, tdarea, vismet)	
63	Eliminate the "fat" boundary cells in any coordinate direction. $mode = 1, 2,$ or 4 indicates the $i, j,$ and k direction, respectively. Set $mode$ equal to the sum of the desired directions. I.e., setting $mode = 5$ will eliminate the "fat" boundary cells in the i and k directions. This test option only applies to structured grids. (mphzmet, nsrhsv, tdarea, tdarea2, vismet)	
64	Remove dt from dq when computing residuals. (12norm1d)	
65	In marching solutions, lower the CFL number for the last marching step. (NSzsolv)	
66	Don't update β in gas3 for equilibrium chemistry. This test option is not recommended but will decrease run time. (gas3)	
67	Flag affecting boundary conditions for one- and two-equation turbulence models at bleed/blowing boundaries. (kebc, sabound, sstbound, TURB_goldbergboundUS, TURB_saboundUS, TURB_sstboundUS, vprtbound, vsctbound)	
	mode Result	
	Treat bleed/blowing boundaries as no-slip walls.	
	1 Treat bleed/blowing boundaries as slip walls. This was the default behavior prior to WIND 5.101.	
	Same as mode 0, except do not use the procedure of Wilcox (2000) to compute the boundary condition for ω in the SST model.	
68	If the density is zero at a coupled boundary, issue a warning, ignore the coupling data, and continue. The default is to issue an error message and abort. This test option only applies to structured grids. (postrbs)	

 Table 15: Non-Production Test Options (Continued)

number	Description
69	By default, (i.e., when this test option is not used) multi-species cases with frozen or finite-rate chemistry solve $ns-1$ species equations, where ns is the number of species, and the mole fraction for species number ns is set so that the total adds to one. All the numerical error is thus concentrated in the value of the mole fraction for the last species.
	When TEST option 69 is set, a "dummy" last species is used, whose mole fraction is always zero. The Fortran variable ns is thus one more than the actual number of species in the flow. We still solve $ns-1$ equations, but that now corresponds to all of the "real" species of interest. In general the total of the mole fractions will not be exactly 1.0, so all the values are rescaled by a constant factor to make it 1.0, spreading out any numerical error over all the species.
	Currently the dummy species is not added automatically. Thus, when this test option is used, the chemistry data (.chm) file must be modified to add entries at the end of the appropriate sections (i.e., thermodynamic coefficients, finite-rate coefficients, and transport properties) for the dummy species. The name of the dummy species doesn't matter, so one method would be to simply copy the entry for one of the other species, and change the species name to DUM to avoid confusion. (bcrcpl, bqlinit, chpstin1, dqlim1, dqlim2, fixer, flagtbad, gas1, gas2, gas3, gas4, propin, qlr2bql, rates, rates2, rates2a, ratesa, ratesaa, ratesadl, ratesadla, ratesarr, ratesbe, ratesbea, ratesf, ratesfa, ratesg, ratesga, stomp, tdespair, transp, updatq1d, updatqnp, US_chemsrc, US_Qcell2Qvrtx)
70	For $mode > 0$, the tolerance level for converging gas properties P , ρ , or T in gas2 will be set to 10^{-mode} . The default is 10^{-4} . This only affects cases using TEST 99. (gas2)

 Table 15: Non-Production Test Options (Continued)

		(********************************
number	Descri	iption
71	equation BC_bcf chpsti Part_E	ation of thermodynamic and transport properties from curve fit ons in .chm file. (aichem, aijkrg, aikeps, airgun, aixyzrg, freebc, BC_bcfreechar, BC_ijkfreechar, BC_usrffreechar, .n1, chrhsv, cpfun, gas5, genturb, gibfn, hfun, kwinflow, Evap, pevap, phinit, pstinp, sfun, tdutv1, therm1, trans1, b, turbupd, uspeci)
	For the	e thermodynamic properties,
	mode	Result
	0	Abort if the temperature exceeds the maximum for the curve fits.
	1	Extrapolate if the temperature exceeds the maximum for the curve fits, and write a warning message to the .lis file.
	2	Same as mode 1, except without a warning message.
	3	If the temperature is outside the range of the curve fits, use the values at the nearest limit.
	4	Use constant values consistent with a gas constant of $287 \text{ m}^2/\text{sec}^2\text{-K}$, and a specific heat at constant pressure of $1004.5 \text{ m}^2/\text{sec}^2\text{-K}$.
	5	If the temperature is outside the range of the curve fits, set the specific heat to its value at the nearest limit and extrapolate for remaining properties.
	Sutherl	e transport properties, the meaning depends on which formula (i.e., land's law or the NASA formula) is being used, and whether the rature is above or below the range of the curve fits.
	mode	Result
	0	For most cases, abort if the temperature is outside the range of the curve fits. The only exception is for Sutherland's law with a temperature lower than the minimum temperature for the curve fits. In that case, the values are extrapolated.
	1	For Sutherland's law, extrapolate if the temperature is outside the range of the curve fits. If the temperature is too high, a warning message is written to the .lis file.
		For the NASA formula, if the temperature is below the minimum value T_{\min} , apply Sutherland's law with constants computed using the NASA formula at T_{\min} . If the temperature is above the maximum value, extrapolate. A warning message is written to the .lis file in both cases.
	2, 4, 5	Same as mode 1, except without a warning message.
	3	If the temperature is outside the range of the curve fits, use the values at the nearest limit.

 Table 15: Non-Production Test Options (Continued)

number	Description
72, 73	For structured grids, periodic boundaries may be established by setting up double (or larger) fringes at the boundaries using GMAN, with the boundary condition type specified as frozen. TEST options 72 and 73 provide the additional information needed to apply the boundary condition.
	TEST option 72 specifies the direction(s) of periodicity, where values of 1, 2, and 4 indicate the i , j , and k direction, respectively. Set $mode$ equal to the sum of the desired directions. E.g., setting $mode = 5$ means periodicity in both the i and k directions.
	TEST option 73 specifies the depth of the fringes.
	As an example, with TEST 72 1 (periodicity in the i direction) and TEST 73 2 (a double fringe), we have
	Conditions at Come from I = 1 I = IDIM - 3 I = 2 I = IDIM - 2 I = IDIM - 1 I = 3 I = IDIM I = 4
	This procedure may only be used for periodic boundaries within a single structured zone, with point-matched grids in the source and target regions. It's primarily useful for cases with higher-order differencing schemes, since the order of the scheme is preserved across the boundary. (kebc, sabound, sstbound, tdbcni, vprtbound, vsctbound)
74	Include k -direction spacing when computing minimum time step in 2-D flows. (tdtmst)
84	Use "old" viscous metric calculation. (emdsolv, vismet)
85	Check for zero volumes when computing viscous metrics. (emdsolv, vismet) $ \frac{mode}{1} \frac{\text{Result}}{\text{Check; if } \leq 0 \text{ print message and continue}} $ $ 2 \text{Check; if } \leq 0 \text{ print message and stop} $
87	Freezes supersonic inflow at initial conditions. (BC_bcfreechar)
88	Bypass checks for negative density and pressure check in tdgas. This is needed for chemically reacting flows if the heat of formation varies widely, since we only have the old heat of formation to use to estimate the temperature. (lpschm, tdgas)
89	Use "old" species flux correction method for multi-species flows. (gas1, transp)

 Table 15: Non-Production Test Options (Continued)

number	Description
90	Chemistry stuff. (chinv) mode Result Analytic chemistry Jacobian (ns = 5 only) Householder chemistry Jacobian Solves chemistry source term explicitly
91	Gas constant. (BC_bcfreebc, chpstin1, therm1) $ \frac{mode}{1} \frac{\text{Result}}{\beta = \gamma = \beta_{\infty}} $ $2 \beta = \gamma = 1.4 $
92	Utilize operator splitting for the reacting chemistry source terms to increase the stability of the integration, allowing more efficient solution of the coupled system. A 4th-order Pade approximation is used to integrate the reaction source terms, with <i>mode</i> setting the number of subiterations. Setting <i>mode</i> = 0 indicates no operator splitting. (chimplicit, chrhss, US_chemsrc, US_chimplicit, US_GaussSeidel)
93	No longer valid; to turn off species diffusion use DIFFUSION NONE in the CHEMISTRY keyword block. (chrhsv, prtinp, rhsvfl, tdutv1, testprtinp)
94	Turn off implicit chemistry terms. This test option only applies to structured grids. (tdimafk, tdimfu, tdimja, tdutaa)
95	Turn off chemistry source term. (rates, rates2, rates2a, ratesa, ratesadl, ratesadla, ratesarr, ratesbe, ratesbea, ratesf, ratesfa, ratesg, ratesga)
96	Gradually apply chemistry source term over the first <i>mode</i> iterations for finite-rate chemistry. (US_chemsrc, chrhss)
97	P. D. Thomas turbulence model scanning direction. By default, Wind-US starts at viscous walls and moves into the field. This test option forces the code to calculate turbulent parameters from any boundary, in addition to walls. (algtur) $ \underbrace{mode}_{0} \underbrace{\text{Result}}_{\text{use } j \text{ lines}} \\ 1 \text{use } k \text{ lines} \\ 2 \text{use } j \text{ and } k \text{ lines} $
99	Initialize finite-rate chemistry with Liu and Vinokur curve fits. (Also see TEST 70.) (gas2) $ \frac{mode}{0} \frac{\text{Result}}{\text{Do not track the species (valid to 50K?)}} $ 1 Track the species (valid to 10K?)

 Table 15: Non-Production Test Options (Continued)

number	Description
100	Time-accurate characteristic extrapolation for coupled, freestream, and inflow boundaries. (BC_bcfree, BC_test100)
	 mode Result Second order, with limit of ΔQ ≤ Q/2 1 1st-order, original characteristic treatment 2 2nd-order, original characteristic treatment (only available for structured grids) 3 1st-order, Roe's average characteristic treatment 4 2nd-order, Roe's average characteristic treatment (only available for structured grids)
102	Use time-averaged back pressure when applying specified mass flow boundary condition at outflow boundaries. (BC_pdsmfr)
104	Treatment of implicit viscous terms. This test option only applies to structured grids. (rcutv1, tdutv1) mode Result Use simple implicit viscous terms. 1 Turn off implicit viscous terms. 2 Use fully-implicit viscous terms. This only applies to the energy diffusion term for frozen and finite-rate chemistry.
105	Time step type. This test option only applies to structured grids. (iterprtinp, tdtmst) $ \frac{mode}{0} \frac{\text{Time Step Type}}{\Delta t = \text{CFL/max}(\lambda_{\xi}, \lambda_{\eta}, \lambda_{\zeta})} $ $1 \text{Flow angle scaling, } \Delta t = \text{CFL} \times (f_{\xi}\Delta \xi + f_{\eta}\Delta \eta + f_{\zeta}\Delta \zeta), \text{ where } f_{\xi} = \sqrt{1 + \tan \theta + \tan \psi} $ $f_{\eta} = f_{\xi} \tan \theta $ $f_{\zeta} = f_{\xi} \tan \psi $ $2 \text{Velocity scaling, } \Delta t = \text{CFL} \times \min(f_{\xi}\Delta \xi, f_{\eta}\Delta \eta, f_{\zeta}\Delta \zeta), \text{ where } f_{\xi} = u/u_{\xi}/ u_{\xi} + c $ $f_{\eta} = u/u_{\eta}/ u_{\eta} + c $ $f_{\zeta} = u/u_{\zeta}/ u_{\zeta} + c $ $3 \Delta t = \text{CFL} \times \min(\Delta \xi, \Delta \eta, \Delta \zeta)/(u + c) $ $4 \Delta t = \text{CFL}/(\lambda_{\xi} + \lambda_{\eta} + \lambda_{\zeta}) $
106	Compute the time step at the start of every cycle (even when Newton time stepping is being used), instead of at the start of every iteration. This test option only applies to structured grids. (axflow, lpgrp, lpmg, redim)

 Table 15: Non-Production Test Options (Continued)

number	Description
107	Flag for choice of Newton and/or second-order time algorithm; 0 to use the NPARC Alliance algorithm, 1 to use the Boeing algorithm. This test option only applies to structured grids. (bbarth, kestep, pstinp, rcimsc, spalart, sst, sst2, sst3, tdimfu, tdimja, tdimsc)
108	Extrapolate freestream outflow. (This only applies to the old GMAN "freestream outflow" boundary type, which is no longer used.) (BC_bcfree, BC_bcfreebc)
109	Boundary flux treatment. This test option only applies to structured grids. (roewal, tdup1) For tdup1:
	$\frac{mode}{0}$ $\frac{\text{Result}}{\text{Conservative}}$ 1 Upwind extrapolation from interior
	For roewal:
	mode Result Characteristic inflow, conservation if flow parallel to wall
	1 Characteristic regardless
	1000 Use conservative wall treatment at all boundaries
110	Grid area variation limiting. This test option only applies to structured grids, and may not be used with third-order fully upwind, fourth-order, or fifth-order explicit operators, specified using the RHS keyword. (roecof)
	$\frac{mode}{Q} = \frac{A_2/A_1}{Q}$
	$egin{array}{ccc} 0 & \infty & & & & & & & & & & & & & & & & &$
	2 1.5
	$3 \qquad 1.33$
	$4 \qquad 1.1$

 Table 15: Non-Production Test Options (Continued)

number	Description
111	Singular matrix check. This test option only applies to structured grids. (jacpr4, jacpr5, jacprg, jacprg4, tdsol4, tdsol9, tdsol11, tdsolg, tdsolg4, tdsolv)
	$\frac{mode}{0}$ $\frac{\text{Result}}{\text{Check}}$; if singular, arbitrarily set diagonal term to 0.1, print a warning, and continue.
	Same as mode 0, but without printing a warning. Don't check.
112	Corrected upwind scheme at boundaries. Defaults to corrected scheme, $mode > 0$ uses second order smoothing with $mode/1000$ as the smoothing level. This test option only applies to structured grids. Users should not use this option. (rhsupw)
113	Check for reverse flow at inflow and outflow boundaries. (BC_bcconf, BC_bcfree)
	 mode
114	Use central differencing in the ζ direction when the Coakley explicit operator is
115	being used, with a central smoothing coefficient of $mode/1000$. (tdup1) Do not rescale inviscid wall total velocity to equal adjacent value, just subtract the normal component from the adjacent value. (BC_bcvel, US_lhswall)
116	Set inward pointing normal to zero in tdbcm1 at unknown grid topology points. This test option only applies to structured grids. (tdbcm1)
117	Freeze conditions at inflow boundaries, even in subsonic flow. (BC_bcfreebc, BC_tst100bc)
118	Singular axis fix. This test option only applies to structured grids. (radavg)
	mode Result 0 Average density, momentum components, and pressure 1 Average density, velocity components, and pressure

 Table 15: Non-Production Test Options (Continued)

number	Description
119	Compute forces and moments using the FOMOCO method (Chan and Buning, 1996). Note that the Wind-US executable must be built with the precompiler directive _FOMOCO defined in order to use this capability; otherwise all the FOMOCO rooutines will be dummies, and will error exit when called. Note also that this requires as input the files <code>mixsur.fmp</code> , <code>grid.ptv</code> , and <code>grid.ibi</code> created by MIXSUR. (lpcycl, lpmg, psrwgv)
121	Under-Relaxation of points adjacent to singular axis. This test option only applies to structured grids. (bcpinw, bcsing, kebc, relsng, sabound, sngthrm, sstbound, vprtbound, vsctbound)
	$\underline{mode} \underline{ ext{Result}}$
	Value on axis is a radius-weighted average of the values at the adjacent points; values at the adjacent points are unchanged
	n Value on axis computed as for mode 0; values at the adjacent points are computed from
	$F_{adj} = (1-r)F_{adj} + rF_{axis}$
	where F_{adj} is the value at the adjacent point, F_{axis} is the axis value, and $r = n/1000$.
122	Allow left-handed coordinates. (tdarea1, tdarea2)
123	Track back pressure, mass flow, and integrated total pressure for outflow boundary conditions, writing the values into the .lis file. The back pressure will be in psi, and the integrated total pressure will be the ratio of the boundary value to the freestream value. When the MASS FLOW keyword is used, the mass flow value will actually be the ratio of the computed value to the desired value. For the other outflow boundary conditions the dimensional mass flow is written, in $\rm lb_m/sec.$
	Note that when the MASS FLOW keyword is used, the back pressure and the mass flow are automatically written into the .lis file, whether this test option is used or not. Specifying TEST 123 will add the integrated total pressure.
	Also note that when the MASS FLOW keyword is used, all three values may be extracted from the .lis file using the resplt (or resplt.pl) utility. For the other outflow boundary conditions, only the integrated total pressure can be extracted. (BC_IntgrtBnd, BC_pdsmfr)
124	Write the accumulated time to the list output (.lis) file every iteration instead of every cycle. This test option only applies to single-zone structured grids, with a CFL number specified using the CFL# keyword mode 1 or 2, and with the trailing "ityp" input value set to 1 to indicate that a global time step should be used equal to the minimum value in the zone. (print_res)
126	Compressibility correction to Baldwin-Lomax turbulence model. (blomax)

 Table 15: Non-Production Test Options (Continued)

number	Description
127 128	Scale printed residual by maximum residual over all time steps. (print_res) Check the L2 norm of the residual for convergence instead of the maximum residual. (GRD_MeshDeform, 12norm1d)
131	Reset the time step for j index values $< mode$ equal to the time step at $j = mode$. I.e., $(\Delta t)_{i,j,k} = (\Delta t)_{i,mode,k}$ for $j < mode$. This is intended for use with boundary layers on $j = 1$ walls, to set the time step in the boundary layer to a (larger) "outer" value. (tdtmst)
132	Renormalize, changing from total to static values. The normalizing values in the .cfl file (e.g., the freestream values in the fpar array in the global header) are unchanged. (redim)
134	Order of extrapolation from interior to boundary values used at freestream, arbitrary inflow, and outflow boundaries, and at coupled boundaries using characteristic coupling. (BC_bcptcond)
	 mode / Result First order. 0, 1 Second order, using a minmod limiter. This mode value only applies to structured grids.
135 136	Resets the time step using a weighting function between the ordinary Euler CFL number and a new "viscous CFL number", for convergence acceleration in viscous layers. The viscous CFL number is set to $mode/1000$. Limited testing indicates that a value of $mode = 50$ is stable and increases the time step near the wall by at least an order of magnitude. This test option only applies to structured grids. (tdtmst) Divergence checking criteria and action. Set $mode = n_1 + 10n_2$, where n_1 and
-50	n_2 are defined below. (lpgrp) n_1 Divergence Definition 1 Max residual > 1.0, L2 norm increasing 2 Max residual > 5.0, L2 norm increasing 3 Max residual > 10.0, L2 norm increasing
	and n_2 Action Taken When Diverging Terminate iteration for current cycle 2 Abort run Reduce CFL number by $1/2$ and terminate iteration for current cycle
137	Butt line interpolation region for USERSPEC; smear USERSPEC conditions over $0.001 \times \text{butt}$ line at minimum and maximum butt line. (uspeci)

 Table 15: Non-Production Test Options (Continued)

number	Description
138	Treatment options for boundary cells. This test option only applies to structured grids. (BC_bcwall, chrhsv, mphzmet, nsrhsv, PreWallBC, tdarea, tdarea1, tdarea2, PreWallBC, US_tdbcg, vismet)
	$\underline{mode} \underline{ ext{Result}}$
	≤ 1 Use large cells
	2 Use large cells, central difference Jacobian
	3 Throw out half cell at boundaries
	5 Solve $\partial P/\partial n$ equation at walls
139	Turn on grid-based flux limiting for the Coakley explicit operator, using the limiting turning angle specified with the GRID LIMITER keyword. (tdup1)
140	Use first-order differencing when computing the $\partial(u, v, w)/\partial \xi$ term in vorticity used in turbulence models. This test option only applies to structured grids. (sst3, vortcy)
141	Use second-order differencing for normal derivatives when applying wall boundary conditions (for explicit boundary conditions only). (BC_bcvel, BC_bcwall, US_tdbcg)
	$\underline{mode} \underline{\text{Result}}$
	1 Second order $\partial p/\partial n$, $\partial T/\partial n$, and $\partial u_{tan}/\partial n$
	Second order $\partial p/\partial n$ and $\partial T/\partial n$, but not $\partial u_{tan}/\partial n$
150	Singular axis on symmetry planes. When symmetry plane test fails, zero this component of velocity. This test option only applies to structured grids. (bcsing)
	mode Result
	$\overline{1}$ $u=0$
	v = 0
	$3 \qquad w = 0$
	4 do not zero any component (use average)
	5 v = w = 0
	$6 \qquad u = w = 0$
	7 u = v = 0
151	For a singular axis, the value on the axis is a radius-weighted average of the values two points away from the axis, instead of the values at the adjacent points; values at the adjacent points are set to the average of the axis value and the value two points away from the axis. E.g., for a singular axis at $j=1$, with the k direction "circumferential", the value on the axis is a radius-weighted average of the values at $j=3$, instead of at $j=2$. Then for each k , the value at $j=2$ is set to the average of the values at $j=1$ and $j=3$. This test option only applies to structured grids, and overrides TEST 121. (bcpinw, bcsing, kebc, kerot, radavg, relsng, sabound, sngthrm, sstbound, vprtbound, vsctbound)

 Table 15: Non-Production Test Options (Continued)

number	Description
153	Iteration frequency for updating pressure at outflow boundaries. The default is 5 for all the DOWNSTREAM PRESSURE modes except VARIABLE and UNSTEADY, where the default is 1. The default is also 1 when outflow boundary conditions are specified using the MASS FLOW, DOWNSTREAM MACH, or COMPRESSOR FACE keyword. (BC_IntgrtBnd)
154	When computing values at "undefined" boundary points, and no neighboring non-hole, non-fringe, points are found, average over neighboring fringe points. This test option only applies to structured grids. (bcundef, kebc, sabound, sstbound, vprtbound, vsctbound)
155	For unstructured grids, extrapolate at freestream characteristic boundaries. (BC_bcfree)
157	Options for USERSPEC inflow. (uspeci)
	 mode 1 Result Put USERSPEC inflow at all points in the buttline range. Do not check for above/below vehicle.
	2 Same as mode 1, but also ignore fuselage station check.
158	Write various unstructured grid info to .lis file. (opngrd)
160	Pressure correction factor = $mode/1000$ for specified mass flow boundary condition. The default is a pressure correction factor of 1. (BC_pdsmfr)
162	Boundary layer edge definition for the Cebeci-Smith turbulence model. (cebeci)
	$\begin{array}{c c} \underline{mode} & \underline{\text{Result}} \\ \hline 0 & 1.0\% \text{ change in } U_{total} \text{ between grid points} \\ 1 & 0.995 \ H_t \\ 2 & 0.99 \ U_{total} \\ 3 & 0.9999 \ U_{total} \end{array}$
163	Criteria for defining F_{max} in Baldwin-Lomax model.
	$\begin{array}{ll} \underline{mode} & \underline{\text{Result}} \\ > 0 & \overline{\text{Search outward from wall; use first peak in } F \text{ that is followed by a} \\ & \text{fractional decrease in } F \text{ of } \underline{mode/1000.} \\ < 0 & \overline{\text{Use the max }} F \text{ value between the wall and the } \underline{ mode } \text{'th grid point from the wall} \\ \end{array}$
	The default is use the first peak in F that is followed by a fractional decrease of 70% (i.e., $mode = 700$). (blomax)
164	Iteration interval for updating gas properties and species for $ireal = 2$ (i.e., equilibrium air chemistry). The default value is 1. $(tdgas)$
165	Sets the tolerance for the distance between grid points used to determine a singular direction to $10^{mode/1000}$. The default is a tolerance of 10^{-8} (i.e., $mode = -8000$). This test option applies to structured grids only. (bcsing, direct)

 Table 15: Non-Production Test Options (Continued)

number	Description
168	In a marching solution with the algebraic turbulence models, begin turbulent flow at streamwise station <i>mode</i> . (tdvis1)
170	NASA Ames time step formula. (CFL increases as $1/\sqrt{\Delta y}$ near the wall. Thus, Δt decreases as $\sqrt{\Delta y}$, not Δy as the default.) C_A is scalar coefficient on CFL; i.e., CFL _{wall} $\propto C_A$. This test option only applies to structured grids, and has no effect if TEST 105 mode 1, 2, or 3 is used. (tdtmst)
	$egin{array}{ccc} rac{mode}{1} & rac{C_A}{0.01} \ n & 0.001n \end{array}$
172	Turn off base energy for ireal = 3 (i.e., frozen and finite-rate chemistry). (aichem, aijkrg, aikeps, aixyzrg, BC_bcfreechar, BC_ijkfreechar, BC_test100, BC_usrffreechar, dqlim1, gas1, gas2, gas3, gas4, gas5, gasint, pstinp, redimchem, stomp, tdimfp, tdroe4, tdroe5, US_EnsightGama, US_EnsightMach, uspeci)
174	For the algebraic turbulence models, update the turbulent viscosity every <i>mode</i> iterations. The default is 1. (tdvis1)
175	Boundary conditions at freestream radial outer boundaries. (BC_bcfree, BC_bcfreechar, tdbcgs)
177	Freeze the maximum residual at the value for the first iteration. This test option applies to structured grids only. (lpgrp)
178	For finite-rate chemistry with multi-species blowing (i.e., using BLOW SPECIES_FRACTIONS with BLOW MASS or BLOW VALVE), gradually introduce the species over the first <i>mode</i> iterations. (BC_bcbled)
179	When solidbody rotation is specified at an arbitrary inflow boundary, modify the rotation radius to get a linear swirl profile in r with the zero velocity point not at the specified center of rotation. $mode = 1000r_0$, where r_0 is the radius (from the point x_c, y_c, z_c specified using the SOLIDBODY keyword in the ARBITRARY INFLOW keyword block) for zero velocity. (rotat)
180	For an actuator disk with free vortex modeling (i.e., the TURNING VORTEX keyword in the ACTUATOR keyword block), set the radius of the solidbody rotation core to $mode/1000$ inches. The default is to set a solidbody core radius such that the pressure at $r = 0$ is $p_{\infty}/10$. (rotat1)
182	For Roe zonal coupling, don't modify the boundary states at interior face points for consistency with boundary values. This test option applies to structured grids only. (tdbcgs)

 Table 15: Non-Production Test Options (Continued)

number	Description
185	When restarting a parabolized marching solution for a case with equilibrium chemistry,
	 <u>mode</u> Result Reset all flow field values at unknown planes, including thermodynamic properties (β, Z, heat of formation, and sound speed), equal to those at the plane upstream of first unknown plane. This should only be necessary when using the RESTART keyword and specifying a restart plane upstream of the last completed plane. Reset thermodynamic properties at the last computed plane to the reference conditions.
	This test option applies to structured grids only. (gasint)
187	The factor for suppression of the streamwise pressure gradient in a marching solution is set to $mode/1000$. The default factor is 0.95 (i.e., $mode=950$), and $mode$ values below 800 are not recommended. When separation or strong adverse pressure gradients are causing problems, values between 800 and 900 will really help. (rhsmar, tdimfp)
189	If a first-order upwind explicit operator modified for stretched grids is used (e.g., RHS ROE FIRST PHYSICAL, then TEST 189 1 must also be specified. (numprtinp)
190	Outgoing wave Roe boundary treatment. This test option applies to structured grids only. (pstinp, roecof, roeht, tdbcgs)
	\underline{mode} Result
	0, 1 Use normal Roe boundary treatment (uses boundary point in formulation)
	2 Lower order by one (does not use boundary point in formulation). This option cannot be used with TVD in the same zone, or with a fourth-order central or fifth-order upwind-biased explicit operator.
	3 Use zero-order extrapolation
192	Save metrics in a temporary file. After the first cycle, metrics will be read rather than computed. This eliminates the CPU resources required to re-compute the metrics each cycle, but adds significant I/O to the computation. In the past, on at least some Cray systems, this reduced the CPU time by approximately 40.8 μ sec/(node-cycle). On the more common platforms, however, it is generally faster to re-compute the metrics rather than store them. (lpcycl)
193	Do <i>not</i> stop if a singular line is encountered normal to a wall. This test option applies to structured grids only. (bbdamp, blinit, kepy2)
194	Bypass singular viscous metric check. This test option applies to structured grids only. (emdsolv, mphzmet, vismet)

 Table 15: Non-Production Test Options (Continued)

number	Description
195	When using BLOW SURFACE, print a warning when the flowfield static pressure becomes larger than the plenum total pressure, causing blowing to be turned off at that point. Note that this is a five-line message written for each iteration and each "closed" node, and could cause the .lis file to become very large very quickly. (BC_bcbled)
196	For overlapping grids, print an error message if there are no interior points adjacent to a fringe point. This test option applies to structured grids only. (BC_norot, srfpar, tdbcgs)
197	Roe self-coupling mode. This test option applies to structured grids only. (pstinp)
	 mode / Result / Once per iteration, using bcself Once per cycle, using standard zone coupling
199	Singular axis averaging — average from 1 to $(max - 1)$, not 1 to max . This test option applies to structured grids only. (bcsing, linzero, radavg, relsng, sngthrm)
200	Don't bomb for negative speed of sound in tdroe3A. This test option applies to structured grids only. (tdroe3A)

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